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## (57) Abstract

There is provided a novel vascular endothelial growth factor, herein designated VEGF-X, in addition to the nucleic acid molecule encoding it, a host cell transformed with said vector and compounds which inhibit or enhance angiogenesis. Also provided is the sequence of a CUB domain present in the sequence of VEGF-X which domain itself prevents angiogenesis and which is used to treat diseases associated with inappropriate vascularisation or angiogenesis.

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#### VASCULAR ENDOTHELIAL GROWTH FACTOR-X

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The present invention is concerned with a novel vascular endothelial growth factor (VEGF) herein designated "VEGF-X", and characterisation of the nucleic acid and amino acid sequences of VEGF-X.

#### Introduction

Angiogenesis involves formation and proliferation of new blood vessels, and is an essential physiological process for normal growth and development of tissues in, for example, embryonic development, tissue regeneration and organ and tissue repair.

Angiogenesis also features in the growth of human cancers which require continuous stimulation of blood vessel growth. Abnormal angiogenesis is associated with other diseases such as rheumatoid arthritis psoriasis and diabetic retinopathy.

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Capillary vessels consist of endothelial cells which carry the genetic information necessary to proliferate to form capillary networks. Angiogenic molecules which can initiate this process have previously been

- characterised. A highly selective mitogen for vascular enothelial cells is vascular endothelial growth factor (VEGF) (Ferrara et al., "Vascular Endothelial Growth Factor: Basic Biology and Clinical Implications". Regulation of angiogenesis, by I.D.
- Goldberg and E.M. Rosen 1997 Birkhauser Verlag
  Basle/Switzerland). VEGF is a potent vasoactive
  protein which is comprised of a glycosylated cationic
  46-49 kd dimer having two 24 kd subunits. It is
  inactivated by sulfhydryl reducing agents and is
- resistant to acidic pH and to heating and binds to immobilised heparin.

VEGF-A has four different forms of 121, 165, 189 and 206 amino acids respectively due to alternative splicing. VEGF121 and VEGF165 are soluble and are capable of promoting angiogenesis, whereas VEGF189 and VEGF206 are bound to heparin containing proteoglycans 5 in the cell surface. The temporal and spatial expression of VEGF has been correlated with physiological proliferation of the blood vessels (Gajdusek, C.M., and Carbon, S.J., Cell Physiol., 139:570-579, (1989)); McNeil, P.L., Muthukrishnan, L., 10 Warder, E., D'Amore, P.A., J. Cell. Biol., 109:811-822, (1989)). Its high affinity binding sites are localized only on endothelial cells in tissue sections (Jakeman, L.B., et al., Clin. Invest. 89:244-253 15 (1989)). The growth factor can be isolated from pituitary cells and several tumor cell lines, and has been implicated in some human gliomas (Plate, K.H. Nature 359:845-848, (1992)). The inhibition of VEGF function by anti-VEGF monoclonal antibodies was shown 20 to inhibit tumor growth in immune-deficient mice (Kim, K.J., Nature 362:841-844, (1993)).

VEGF proteins have been described in the following patents and applications all of which are hereby incorporated by reference EP-0,506,477, WO-95/24473, WO-98/28621, WO-90/13649, EP-0,476,983, EP-0,550,296, WO-90/13649, WO-96/26736, WO-96/27007, WO-98/49300, WO-98/36075, WO-98/840124, WO-90/11084, WO-98/24811, WO-98/10071, WO-98/07832, WO-98/02543, WO-97/05250, WO-91/02058, WO-96/39421, WO-96/39515, WO-98/16551.

The present inventors have now identified a further vascular endothelial growth factor, designated herein as "VEGF-X", and the nucleic acid sequence encoding it, which has potentially significant benefits for the treatment of tumours and other conditions mediated by inappropriate angiogenic activity.

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## Summary of the Invention

In the present application, there is provided a novel vascular endothelial growth factor, herein designated "VEGF-X", nucleic acid molecules encoding said growth factor, an expression vector comprising said nucleic acid molecule, a host cell transformed with said vector and compounds which inhibit or enhance angiogenesis. Also provided is the sequence of a CUB domain present in the sequence of VEGF-X which domain itself prevents angiogenesis and which is used to treat diseases associated with inappropriate vascularisation or angiogenesis.

## 15 <u>Detailed Description of the Invention</u>

Therefore, according to a first aspect of the present invention there is provided a nucleic acid molecule encoding a VEGF-X protein or a functional equivalent, 20 fragment, derivative or bioprecursor thereof, said protein comprising the amino acid sequence from position 23 to 345 of the amino acid sequence illustrated in Figure 10. Alternatively, the nucleic acid molecule of the invention encodes the complete 25 sequence identified in Figure 10 and which advantageously includes a signal peptide to express said protein extracellularly. Preferably, the nucleic acid molecule is a DNA and even more preferably a cDNA molecule. Preferably, the nucleic acid molecule 30 comprises the nucleotide sequence from position 257 to 1291 of the nucleotide sequence illustrated in Figure In a preferred embodiment the nucleic acid is of mammalian origin and even more preferably of human origin.

In accordance with the present invention a functional

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equivalent should be taken to mean a protein, or a sequence of amino acids that have similar function to the VEGF-X protein of the invention.

Also provided by this aspect of the present invention is a nucleic acid molecule such as an antisense molecule capable of hybridising to the nucleic acid molecules according to the invention under high stringency conditions, which conditions would be well known to those skilled in the art.

Stringency of hybridisation as used herein refers to conditions under which polynucleic acids are stable. The stability of hybrids is reflected in the melting temperature (Tm) of the hybrids. Tm can be approximated by the formula:

## $81.5^{\circ}C+16.6(\log_{10}[Na^{+}]+0.41 (\$G&C)-600/1$

wherein 1 is the length of the hybrids in nucleotides. Tm decreases approximately by 1-1.5°C with every 1% decrease in sequence homology.

The term "stringency" refers to the hybridisation conditions wherein a single-stranded nucleic acid joins with a complementary strand when the purine or pyrimidine bases therein pair with their corresponding base by hydrogen bonding. High stringency conditions favour homologous base pairing whereas low stringency conditions favour non-homologous base pairing.

"Low stringency" conditions comprise, for example, a temperature of about 37°C or less, a formamide concentration of less than about 50%, and a moderate to low salt (SSC) concentration; or, alternatively, a temperature of about 50°C or less, and a moderate to high salt (SSPE) concentration, for example 1M NaCl.

"High stringency" conditions comprise, for example, a temperature of about 42°C or less, a formamide concentration of less than about 20%, and a low salt (SSC) concentration; or, alternatively, a temperature of about 65°C, or less, and a low salt (SSPE) concentration. For example, high stringency conditions comprise hybridization in 0.5 M NaHPO4, 7% sodium dodecyl sulfate (SDS), 1 mM EDTA at 65°C (Ausubel, F.M. et al. Current Protocols in Molecular Biology, Vol. I, 1989; Green Inc. New York, at 2.10.3).

"SSC" comprises a hybridization and wash solution. A stock 20X SSC solution contains 3M sodium chloride, 0.3M sodium citrate, pH 7.0.

"SSPE" comprises a hybridization and wash solution. A 1X SSPE solution contains 180 mM NaCl, 9mM Na<sub>2</sub>HPO<sub>4</sub> and 1 mM EDTA, pH 7.4.

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The nucleic acid capable of hybridising to nucleic acid molecules according to the invention will generally be at least 70%, preferably at least 80 or 90% and more preferably at least 95% homologous to the nucleotide sequences according to the invention.

The antisense molecule capable of hybridising to the nucleic acid according to the invention may be used as a probe or as a medicament or may be included in a pharmaceutical composition with a pharmaceutically acceptable carrier, diluent or excipient therefor.

The term "homologous" describes the relationship between different nucleic acid molecules or amino acid sequences wherein said sequences or molecules are related by partial identity or similarity at one or more blocks or regions within said molecules or

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sequences.

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The present invention also comprises within its scope proteins or polypeptides encoded by the nucleic acid molecules according to the invention or a functional equivalent, derivative or bioprecursor thereof.

Therefore, according to a further aspect of the present invention, there is provided a VEGF-X protein, or a functional equivalent, derivative or bioprecursor 10 thereof, comprising an amino acid sequence from position 23 to 345 of the sequence as illustrated in Figure 10, or alternatively which amino acid sequence comprises the complete sequence of Figure 10. A further aspect of the invention comprises a VEGF-X 15 protein, or a functional equivalent, derivative or bioprecusor thereof, encoded by a nucleic acid molecule according to the invention. Preferably, the VEGF-X protein encoded by said nucleic acid molecule comprises the sequence from position 23 to 345 of the 20 amino acid sequence as illustrated in Figure 10, or which sequence alternatively comprises the sequence of amino acids of Figure 10.

The DNA molecules according to the invention may, advantageously, be included in a suitable expression vector to express VEGF-X encoded therefrom in a suitable host. Incorporation of cloned DNA into a suitable expression vector for subsequent transformation of said cell and subsequent selection of the transformed cells is well known to those skilled in the art as provided in Sambrook et al. (1989), molecular cloning, a laboratory manual, Cold Spring Harbour Laboratory Press.

An expression vector according to the invention includes a vector having a nucleic acid according to

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the invention operably linked to regulatory sequences, such as promoter regions, that are capable of effecting expression of said DNA fragments. The term "operably linked" refers to a juxta position wherein the components described are in a relationship 5 permitting them to function in their intended manner. Such vectors may be transformed into a suitable host cell to provide for expression of a polypeptide according to the invention. Thus, in a further aspect, the invention provides a process for preparing 10 polypeptides according to the invention which comprises cultivating a host cell, transformed or transfected with an expression vector as described above under conditions to provide for expression by the vector of a coding sequence encoding the 15 polypeptides, and recovering the expressed polypeptides.

The vectors may be, for example, plasmid, virus or phage vectors provided with an origin of replication, and optionally a promoter for the expression of said nucleotide and optionally a regulator of the promoter.

The vectors may contain one or more selectable

25 markers, such as, for example, ampicillin resistance.

Regulatory elements required for expression include promoter sequences to bind RNA polymerase and transcription initiation sequences for ribosome binding. For example, a bacterial expression vector may include a promoter such as the lac promoter and for translation initiation the Shine-Dalgarno sequence and the start codon AUG. Similarly, a eukaryotic expression vector may include a heterologous or homologous promoter for RNA polymerase II, a downstream polyadenylation signal, the start codon AUG, and a termination codon for detachment of the

ribosome. Such vectors may be obtained commercially or assembled from the sequences described by methods well known in the art.

Nucleic acid molecules according to the invention may 5 be inserted into the vectors described in an antisense orientation in order to provide for the production of antisense RNA. Antisense RNA or other antisense nucleic acids may be produced by synthetic means.

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In accordance with the present invention, a defined nucleic acid includes not only the identical nucleic acid but also any minor base variations including in particular, substitutions in cases which result in a synonymous codon (a different codon specifying the same amino acid residue) due to the degenerate code in conservative amino acid substitutions. The term "nucleic acid sequence" also includes the complementary sequence to any single stranded sequence given regarding base variations.

The present invention also advantageously provides nucleic acid sequences of at least approximately 10 contiguous nucleotides of a nucleic acid according to the invention and preferably from 10 to 50 nucleotides even more preferably, the nucleic acid sequence comprise the sequences illustrated in Figure 3. These sequences may, advantageously be used as probes or primers to initiate replication, or the like. Such nucleic acid sequences may be produced according to techniques well known in the art, such as by recombinant or synthetic means. They may also be used in diagnostic kits or the like for detecting the presence of a nucleic acid according to the invention. These tests generally comprise contacting the probe with the sample under hybridising conditions and detecting for the presence of any duplex or triplex

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formation between the probe and any nucleic acid in the sample.

The nucleic acid sequences according to this aspect of the present invention comprise the sequences of nucleotides illustrated in Figures 3 and 5.

According to the present invention these probes may be anchored to a solid support. Preferably, they are present on an array so that multiple probes can simultaneously hybridize to a single biological sample. The probes can be spotted onto the array or synthesised in situ on the array. (See Lockhart et al., Nature Biotechnology, vol. 14, December 1996 "Expression monitoring by hybridisation to high density oligonucleotide arrays". A single array can contain more than 100, 500 or even 1,000 different probes in discrete locations.

The nucleic acid sequences, according to the invention 20 may be produced using such recombinant or synthetic means, such as for example using PCR cloning mechanisms which generally involve making a pair of primers, which may be from approximately 10 to 50 25 nucleotides to a region of the gene which is desired to be cloned, bringing the primers into contact with mRNA, cDNA, or genomic DNA from a human cell, performing a polymerase chain reaction under conditions which brings about amplification of the 30 desired region, isolating the amplified region or fragment and recovering the amplified DNA. Generally, such techniques are well known in the art, such as described in Sambrook et al. (Molecular Cloning: a Laboratory Manual, 1989).

The nucleic acids or oligonucleotides according to the invention may carry a revealing label. Suitable

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labels include radioisotopes such as <sup>32</sup>P or <sup>35</sup>S, enzyme labels or other protein labels such as biotin or fluorescent markers. Such labels may be added to the nucleic acids or oligonucleotides of the invention and may be detected using known techniques per se.

Advantageously, human allelic variants or polymorphisms of the DNA molecule according to the invention may be identified by, for example, probing cDNA or genomic libraries from a range of individuals, for example, from different populations. Furthermore, nucleic acids and probes according to the invention may be used to sequence genomic DNA from patients using techniques well known in the art, such as the Sanger Dideoxy chain termination method, which may, advantageously, ascertain any predisposition of a patient to certain disorders associated with a growth factor according to the invention.

20 The protein according to the invention includes all possible amino acid variants encoded by the nucleic acid molecule according to the invention including a polypeptide encoded by said molecule and having conservative amino acid changes. Conservative amino 25 acid substitution refers to a replacement of one or more amino acids in a protein as identified in Table 1. Proteins or polypeptides according to the invention further include variants of such sequences, including naturally occurring allelic variants which are substantially homologous to said proteins or 30 polypeptides. In this context, substantial homology is regarded as a sequence which has at least 70%, preferably 80 or 90% and preferably 95% amino acid homology with the proteins or polypeptides encoded by the nucleic acid molecules according to the invention. 35 The protein according to the invention may be recombinant, synthetic or naturally occurring, but is

preferably recombinant.

The nucleic acid or protein according to the invention may be used as a medicament or in the preparation of a medicament for treating cancer or other diseases or conditions associated with expression of VEGF-X protein.

Advantageously, the nucleic acid molecule or the
protein according to the invention may be provided in
a pharmaceutical composition together with a
pharmacologically acceptable carrier, diluent or
excipient therefor.

The present invention is further directed to 15 inhibiting VEGF-X in vivo by the use of antisense technology. Antisense technology can be used to control gene expression through triple-helix formation of antisense DNA or RNA, both of which methods are based on binding of a polynucleotide to DNA or RNA. 20 For example, the 5' coding portion or the mature DNA sequence, which encodes for the protein of the present invention, is used to design an antisense RNA oligonucleotide of from 10 to 50 base pairs in length. A DNA oligonucleotide is designed to be complementary 25 to a region of the gene involved in transcription (triple-helix - see Lee et al. Nucl. Acids Res., 6:3073 (1979); Cooney et al., Science, 241:456 (1988); and Dervan et al., Science, 251: 1360 (1991), thereby preventing transcription and the production of VEGF-X. 30 The antisense RNA oligonucleotide hybridises to the mRNA in vivo and blocks translation of an mRNA molecule into the VEGF-X protein (antisense - Okano, J. Neurochem., 56:560 (1991); Oligodeoxynucleotides as Antisense Inhibitors of Gene Expression, CRC Press, 35 Boca Raton, FL (1988)).

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Alternatively, the oligonucleotide described above can be delivered to cells by procedures in the art such that the anti-sense RNA and DNA may be expressed in vivo to inhibit production of VEGF-X in the manner described above.

Antisense constructs to VEGF-X, therefore, may inhibit the angiogenic activity of VEGF-X and prevent the further growth of or even regress solid tumours, since angiogenesis and neovascularization are essential steps in solid tumour growth. These antisense constructs may also be used to treat rheumatoid arthritis, psoriasis and diabetic retinopathy which are all characterized by abnormal angiogenesis.

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A further aspect of the invention provides a host cell or organism, transformed or transfected with an expression vector according to the invention. The host cell or organism may advantageously be used in a method of producing VEGF-X, which comprises recovering any expressed VEGF-X from the host or organism transformed or transfected with the expression vector.

According to a further aspect of the invention there 25 is also provided a transgenic cell, tissue or organism comprising a transgene capable of expressing VEGF-X protein according to the invention. The term "transgene capable of expression" as used herein means a suitable nucleic acid sequence which leads to 30 expression of VEGF-X or proteins having the same function and/or activity. The transgene, may include, for example, genomic nucleic acid isolated from human cells or synthetic nucleic acid, including DNA integrated into the genome or in an extrachromosomal 35 state. Preferably, the transgene comprises the nucleic acid sequence encoding the proteins according to the invention as described herein, or a functional

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fragment of said nucleic acid. A functional fragment of said nucleic acid should be taken to mean a fragment of the gene comprising said nucleic acid coding for the proteins according to the invention or a functional equivalent, derivative or a nonfunctional derivative such as a dominant negative mutant, or bioprecursor of said proteins. For example, it would be readily apparent to persons skilled in the art that nucleotide substitutions or deletions may be used using routine techniques, which do not affect the protein sequence encoded by said nucleic acid, or which encode a functional protein according to the invention.

- 15 VEGF-X protein expressed by said transgenic cell, tissue or organism or a functional equivalent or bioprecursor of said protein also forms part of the present invention.
- 20 Antibodies to the protein or polypeptide of the present invention may, advantageously, be prepared by techniques which are known in the art. For example, polyclonal antibodies may be prepared by inoculating a host animal, such as a mouse or rabbit, with the 25 polypeptide according to the invention or an epitope thereof and recovering immune serum. Monoclonal antibodies may be prepared according to known techniques such as described by Kohler R. and Milstein C., Nature (1975) 256, 495-497. Advantageously, such 30 antibodies may be included in a kit for identifying VEGF-X in a sample, together with means for contacting the antibody with the sample.

Advantageously, the antibody according to the invention may also be used as a medicament or in the preparation of a medicament for treating tumours or other diseases associated with expression of VEGF-X.

The invention also further provides a pharmaceutical composition comprising said antibody together with a pharmaceutically acceptable carrier diluent or excipient therefor.

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Proteins which interact with the polypeptide of the invention may be identified by investigating protein-interactions using the two-hybrid vector system first proposed by Chien et al., (1991) Proc. Natl. Acad.

10 Sci. USA 88 : 9578-9582.

This technique is based on functional reconstitution in vivo of a transcription factor which activates a reporter gene. More particularly the technique comprises providing an appropriate host cell with a DNA construct comprising a reporter gene under the control of a promoter regulated by a transcription factor having a DNA binding domain and an activating domain, expressing in the host cell a first hybrid DNA sequence encoding a first fusion of a fragment or all of a nucleic acid sequence according to the invention and either said DNA binding domain or said activating domain of the transcription factor, expressing in the host at least one second hybrid DNA sequence, such as a library or the like, encoding putative binding proteins to be investigated together with the DNA binding or activating domain of the transcription factor which is not incorporated in the first fusion; detecting any binding of the proteins to be investigated with a protein according to the invention by detecting for the presence of any reporter gene product in the host cell; optionally isolating second hybrid DNA sequences encoding the binding protein.

An example of such a technique utilises the GAL4 protein in yeast. GAL4 is a transcriptional activator of galactose metabolism in yeast and has a separate

domain for binding to activators upstream of the galactose metabolising genes as well as a protein binding domain. Nucleotide vectors may be constructed, one of which comprises the nucleotide residues encoding the DNA binding domain of GAL4. 5 These binding domain residues may be fused to a known protein encoding sequence, such as for example, the nucleic acids according to the invention. The other vector comprises the residues encoding the protein binding domain of GAL4. These residues are fused to 10 residues encoding a test protein. Any interaction between polypeptides encoded by the nucleic acid according to the invention and the protein to be tested leads to transcriptional activation of a reporter molecule in a GAL-4 transcription deficient 15 yeast cell into which the vectors have been transformed. Preferably, a reporter molecule such as β-galactosidase is activated upon restoration of transcription of the yeast galactose metabolism genes.

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A further aspect of the present invention also provides a method of identifying VEGF-X in a sample, which method comprises contacting said sample with an antibody according to the invention and monitoring for any binding of any proteins to said antibody. A kit for identifying the presence of VEGF-X in a sample is also provided comprising an antibody according to the invention and means for contacting said antibody with said sample.

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VEGF-X may be recovered and purified from recombinant cell cultures by methods known in the art, including ammonium sulfate or ethanol precipitation, acid extraction, anion or cation exchange chromatography, phosphocellulose chromatography, hydrophobic interaction chromatography, affinity chromatography, hydroxyapatite chromatography and lectin

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chromatography.

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The VEGF-X protein of the present invention may be a naturally purified product, or a product of chemical synthetic procedures, or produced by recombinant techniques from a prokaryotic or eukaryotic host (for example, by bacterial yeast, higher plant, insect and mammalian cells in culture). Depending upon the host employed in a recombinant production procedure, the polypeptides of the present invention may be glycosylated with mammalian or other eukaryotic carbohydrates or may be non-glycosylated.

VEGF-X is particularly advantageous as a wound healing 15 agent, where, for example, it is necessary to revascularize damaged tissues, or where new capillary angiogenesis is important. Accordingly, VEGF-X may be used for treatment of various types of wounds such as for example, dermal ulcers, including pressure sores, 20 venous ulcers, and diabetic ulcers. In addition, it can be used in the treatment of full-thickness burns and injuries where angiogenesis is desired to prepare the burn in injured sites for a skin graft and flap. In this case, VEGF-X or the nucleic acid encoding it may be applied directly to the wound. VEGF-X may be 25 used in plastic surgery when reconstruction is required following a burn, other trauma, or even for cosmetic purposes.

An important application of VEGF-X is to induce the growth of damaged bone, periodontium or ligament tissue. For example, it may be used in periodontal disease where VEGF-X is applied to the roots of the diseased teeth, leading to the formation of new bone and cementum with collagen fibre ingrowths. It can be used for regenerating supporting tissues of teeth, including alveolar bone, cementum and periodontal

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ligament, that have been damaged by disease and trauma.

Since angiogenesis is important in keeping wounds clean and non-infected, VEGF-X may be used in association with surgery and following the repair of It should be particularly useful in the treatment of abdominal wounds where there is a high risk of infection.

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VEGF-X can also be used for the promotion of endothelialization in vascular graft surgery. In the case of vascular grafts using either transplanted or synthetic material, VEGF-X may be applied to the surface of the graft or at the junction to promote the growth of the vascular endothelial cells. One derivation of this is that VEGF-X can be used to repair the damage of myocardial and other occasions where coronary bypass surgery is needed by stimulating the growth of the transplanted tissue. Related to this is the use of VEGFX to repair the cardiac vascular system after ischemia.

The protein of the present invention may also be employed in accordance with the present invention by expression of such protein in vivo, which is often referred to as "gene therapy".

Thus, for example, cells such as bone marrow cells may be engineered with a polynucleotide (DNA or RNA) 30 encoding for the protein ex vivo as defined herein, the engineered cells are then provided to a patient to be treated with the polypeptide. Such methods are well-known in the art. For example, cells may be engineered by procedures known in the art by use of a 35 retroviral particle containing RNA encoding for the protein of the present invention.

Similarly, cells may be engineered in vivo for expression of the protein in vivo, for example, by procedures known in the art.

A further aspect of the invention comprises a method of treating a disorder mediated by expression of a protein according to the invention, by administering to a patient an amount of an antisense molecule as described herein, in sufficient concentration to alleviate or reduce the symptoms of said disorder.

Compounds which inhibit or enhance angiogenesis may be identified by providing a host cell or organism according to the invention or a transgenic cell, tissue or organism according to the invention, contacting a test compound with said cell, tissue or organism and monitoring for the effect of said compound compared to a cell tissue or organism which has not been contacted with said compound. These compounds may themselves be used as a medicament or included in a pharmaceutical composition for treatment of disorders mediated by inappropriate vascularisation or angiogenic activity.

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The present inventors have also, advantageously, identified in the sequence encoding the VEGF-X protein a CUB domain, which has heretofore not previously been identified in VEGF-type growth factors. The VEGF-X protein may therefore exert dual regulatory effects via interaction with the VEGF tyrosine kinase receptors or with neuropilin receptors mediated by the CUB domain. Thus, the sequence encoding said CUB domain may be included in an expression vector for subsequent transformation of a host cell, tissue or organism.

VEGF-X or fragments thereof may be able to modulate

the effects of pro-angiogenic growth factors such as VEGF as indicated in the findings presented in the examples below that the N-terminal part of the VEGF-X protein, a CUB-like domain, is able to inhibit VEGF-stimulated proliferation of HUVECs. VEGF-X or fragments thereof may therefore be useful in therapy of conditions involving inappropriate angiogenesis. Inhibition of the angiogenic activity of VEGF has been linked with inhibition of tumour growth in several models eg Kim K. J. et al, Nature 362:841-844, (1993). Additionally, agents able to inhibit angiogenesis would be expected to be useful in treating other angiogenesis-dependent diseases such a retinopathy, osteoarthritis and psoriasis(Folkman, J., Nature Medicine 1:27-31, (1995).

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As identified in more detail in the Examples described herein the present inventors have surprisingly identified that the CUB domain of VEGF-X is able to inhibit stimulation of proliferation of HUVECs induced by either VEGF or bFGF. The CUB domain may, therefore, be utilised as a therapuetic agent for inhibition of angiogenesis and for treatment of condition associated with inappropriate vascularisation or angiogenesis.

Therefore according to a further aspect of the invention there is provided a method of inhibiting angiogenic activity and inappropriate vascularisation including formation and proliferation of new blood vessels, growth and development of tissues, tissue regeneration and organ and tissue repair in a subject said method comprising administering to said subject an amount of a polypeptide having an amino acid sequence from position 40 to 150 of the sequence illustrated in Figure 10 or a nucleic acid molecule encoding the CUB domain according to the invention in

- 20 -

sufficient concentration to reduce or prevent said angiogenic activity.

Furthermore there is also provided a method of treating or preventing any of cancer, rheumatoid arthritis, psoriasis and diabetic retinopathy, said method comprising administering to said subject an amount of a polypeptide having an amino acid sequence from position 40 to 150 of the sequence illustrated in Figure 10 or a nucleic acid molecule encoding the CUB domain according to the invention in sufficient concentration to treat or prevent said disorders.

The CUB domain may also be used to identify compounds that inhibit or enhance angiogenic activity such as 15 inappropriate vascularisation, in a method comprising contacting a cell expressing a VEGF receptor and/or a neuropilin 1 or 2 type receptor with said compound in the presence of a VEGF-X protein according to the invention and monitoring for the effect of said 20 compound or said cell when compared to a cell which has not been contacted with said compound. Such compounds may then be used as appropriate to prevent or inhibit angiogenic activity to treat the disorders or conditions described herein, or in a 25 pharmaceutical composition. An antibody to said CUB domain may also be useful in identifying other proteins having said sequences.

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## Deposited Plasmids

	Plasmid VEGFX/p	Date of Deposit	Accession No.
5	1TOPO FL	1 March 1999	LMBP 3925
10	Plasmid VEGFX/p amino acids G230-G345	DRSETB BD 1 March 1999	LMBP 3926
	Plasmid VEGFX/p	20 October 1999	LMBP 3977
15	Plasmid VEGF-X PET22b	CUB 20 December 1999	
20	Coordinated Co.	nids were deposited at th llections of Microorganis oor Moleculaire Biologie-	sms (BCCM) at
20	Plasmidencolle	ctie (LMBP) B-9000, Ghent h the provisions of the E	, Belgium, in
25	reference to the	may be more clearly under he accompanying example, ry, with reference to the ein:	which is
30	Figure 1:	is a DNA sequence identificate LifeSeq <sup>TM</sup> database novel VEGF-X protein.	
35	Figure 2:	is an illustration of ar sequence of the nucleic of Figure 1.	

sequences utilised to identify the VEGF-X protein according to the invention.  Figure 4: is a diagrammatic illustration of the spatial relationships in the VEGF-X sequence of the clones identified using the PCR primer sequences of Figure 3.  Figure 5: is an illustration of the nucleotide sequences of the 5' RACE primers used to identify the 5' end of the VEGF-X open reading frame.  Figure 6: is an illustration of the sequence obtained from the RACE experiment.  Pigure 7: is an illustration of the nucleotide sequences obtained from the search of LifeSeqTM database using the sequence in Figure 6.  Figure 8: is an illustration of the primers used to clone the entire coding sequence of VEGF-X.  Figure 9: is an illustration of the entire coding sequence of Figure 9: is an illustration of the predicted amino acid sequence of the nucleotide sequence of Figure 9.  Figure 11: is an alignment of the sequence of		Figure 3:	is an illustration of PCR primer
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sequence of Figure 9.			
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Figure 11: is an alignment of the sequence of	35		
		Figure 11:	is an alignment of the sequence of

Figure 10 with the sequences of VEGF-A to D.

protein according to the invention.

is an illustration of variant Figure 12: sequences of the VEGF-X protein 5 according to the invention. is an illustration of the Figure 13: oligonucleotide primers used for E.coli expression of VEGF-X domains 10 and for expression of the full length sequence of VEGF-X in a baculovirus/insect cell expression system. 15 depicts nucleic acid sequences of 18 Figure 14: human EST clones obtained from a BLAST search of the LifeSeq $^{\text{TM}}$  database used to identify the full sequence encoding VEGF-X. 20 depicts the nucleotide sequences of 50 Figure 15: human EST clones obtained from the LifeSeqTM database. 25 is an illustration of nucleotide Figure 16: sequences utilised as primers to identify the nucleotide sequence encoding VEGF-X. 30 is a nucleotide sequence coding for a Figure 17: partial VEGF-X protein according to the invention. is an illustration of a partial Figure 18: 35 nucleotide sequence encoding VEGF-X

5	Figure 19:	is an illustration of a DNA and polypeptide sequence used for mammalian cell expression of VEGF-X. The predicted VEGF-X signal sequence is in lower case letters. The C-terminal V5 epitope and His6 sequences are underlined.
10	Figure 20:	is an illustration of a DNA and polypeptide sequence used for baculovirus/insect cell expression of VEGF-X. In the polypeptide sequence
15		the signal sequence is shown in lower case. The N-terminal peptide tag added to the predicted mature VEGF-X sequence is underlined.
20	Figure 21:	is an illustration of a DNA and polypeptide sequence used for <i>E. coli</i> expression of VEGF-X. The polypeptide sequences at the N- and C- termini derived from the MBP fusion and His6 tag respectively are underlined.
25	Figure 22:	illustrates the disulphide-linked dimerisation of VEGF-X. Protein samples were analysed by SDS-PAGE. Prior to loading the gel, samples were heated to 95°C for 5 minutes in sample
30		buffer in the presence (+) or absence (-) of reducing agent. (A) samples from COS cell expression of a C- terminally V5/His6 peptide-tagged construct. The left hand panel is
35		total conditioned medium, the right hand panel is material purified on Nickel agarose resin. Reduced monomer

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and putative disulphide-linked, nonreduced dimer are indicated by arrows. There appears to be proteolysis of the protein during purification. Gels were 5 blotted onto nylon membranes and protein detected with an anti V5 monoclonal antibody. (B) Samples from E.coli expression of a maltose-binding protein/His6 dual fusion construct. M 10 indicates the molecular weight markers (Benchmark, LifeTechnologies). gel was stained with Coomassie Blue by standard procedures. The fusion protein has an apparent molecular 15 weight of 80kDa. Figure 23: illustrates the glycosylation of VEGF-X. VEGF-X was purified from the culture supernatant of COS cells 20 transfected with the pcDNA6/V5-His construct. Supernatants were harvested 72h post-transfection and purified on nickel resin. Samples were then treated with EndoH (+) or 25 untreated (-) before SDS-PAGE and blotting, as described in the legend to Figure 22. Figure 24: is an illustration of the DNA and 30 polypeptide sequence used for E. coli expression of the VEGF-like domain of VEGF-X. Polypeptide sequences at the N-terminus of the protein derived from the vector are underlined. 35 Figure 25: shows expression of the VEGF-X VEGF

domain in E. coli. Lane 1-10µl broad

range marker (New England Biolabs), lane 2-10µl unreduced sample, lane 3-10µl reduced sample. The reduced PDGF domain protein (lane 3) has an 5 apparent molecular weight of approximately 19kDa on SDS-PAGE. Figure 26: illustrates a DNA and polypeptide sequence used for E. coli expression 10 of the CUB-like domain of VEGF-X. polypeptide sequence at the N-terminus derived from the vector-encoded signal and the introduced His6 tag are underlined. 15 shows expression of the VEGF-X CUB Figure 27: domain in E. coli. The CUB domain protein was purified on Nickel chelate resin. The protein migrates at 20 approximately 23kDa on SDS-PAGE. Figure 28: illustrates the effect of truncated VEGF-X (CUB domain) on HUVEC proliferation. (A) Human Umbilical 25 Vein Endothelial Cells (one-daytreatment). (B) Human Umbilical Vein Endothelial Cells (24-hour starving followed by one-day-treatment). (C) Effect of VEGF-A165 and VEGF-X CUB 30 domain on the proliferation of HUVEC (two-day-treatment). Figure 29: depicts the tissue distribution of VEGF-X mRNA analysed by Northern 35 blotting and RT-PCR in (A) normal tissues and (B) tumour tissue and cell lines.

	Figure 30:	depicts the partial intron/exon
		structure of the VEGF-X gene. (A)
		Genomic DNA sequences of 2 exons
		determined by sequencing; exon
5		sequence is in upper case, intron
		sequence is in lower case. (B) Shows
		the location of splice sites within
		the VEGF-X cDNA sequence. The
		location of mRNA splicing events is
10		indicated by vertical lines. The
		cryptic splice donor/acceptor site at
	,	nt. 998/999 (diagonal lines) gives
		rise to the splice variant forms of
		VEGF-X. No splice site information is
15		given for the region shown in italics.
	Pilanona 21	
	Figure 31:	is a graphic representation of the
		effect of FL-VEGF-X on HuVEC
20		proliferation: (24 hour serum
		starvation followed by one day treatment).
		or daring to
	Figure 32:	is a graphic representation of the
		combined effect of truncated VEGF-X
25		(CUB domain) and human recombinant
		VEGF <sub>165</sub> on HuVEC proliferation: (24 hour
		serum starvation followed by two day
		treatment).
30	Figure 33.	
30	Figure 33:	is a graphic representation of the
		combined effect of the CUB domain and human recombinant bFGF on HuVEC
	•	proliferation: (24 hour serum
٠		starvation followed by two day
35	·	treatment).
		-
	Figure 34:	is a graphic representation of the

results of a LDH assay for testing cytotoxicity of the CUB domain or the CUB domain with rhVEGF<sub>165</sub>.

- 5 Figure 35: is a graphic representation of the results obtained from a LDH assay for testing cytotoxicity of the CUB domain or CUB domain with rh-bFGF.
- 10 A BLAST (Basic Local Alignment Search Tool; Altschul et al., 1990 J. Mol. Biol. 215, 403-410) search was performed in the proprietary LifeSeqTM human EST database (Incyte Pharmaceuticals, Inc., Palo Alto, CA, USA). BLAST produces alignments of both nucleotide and amino acid sequences to determine 15 sequence similarity. Because of the local nature of the alignments, BLAST is especially useful in determining exact matches or in identifying homologues. While it is useful for matches which do 20 not contain gaps, it is inappropriate for performing motif-style searching. The fundamental unit of BLAST algorithm output is the High-scoring Segment Pair (HSP).
- 25 Eighteen human EST clones (Figure 14) with high similarity to the previously identified VEGF proteins were identified and a further fifty EST clones (Figure 15) were identified using these sequences as query sequences, allowing us to deduce the putative 30 sequence for the new VEGF-X protein. The sequences obtained were compared to known sequences to determine regions of homology and to identify the sequence as a novel VEGF-type protein. Using the DNA sequence information in the databases we were able to 35 prepare suitable primers having the sequences of VEGF-X 1-10 illustrated in Figure 3 for use in subsequent RACE experiments to obtain the complete

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DNA sequence for the VEGF-X gene.

## Cloning

A profile was developed based on the VEGF-like domain in existing VEGF sequences (VEGF-A, B, C and D). This was used to search the public databases and the Incyte LifeSeq<sup>TM</sup> database. No significant novel matching sequences were found in the public databases. All of the matching sequences found in

the LifeSeq<sup>TM</sup> database (~1000) were assembled to give a smaller number of sequences (~30), which included the known VEGFs and a potential novel VEGF (figures 1 and 2). This sequence was named VEGF-X.

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Oligonucleotides were designed to amplify the VEGF-X sequence from cDNA (figure 3). The ESTs found in LifeSeq<sup>TM</sup> were from a range of tissues, with a slight predominance of sequences from ovary, testis,

- placenta and lung (Figure 14 and 15). Accordingly the oligonucleotides were used to amplify cDNA derived from lung and placenta. First-round PCR products were found at ~200bp larger than the expected sizes, while 3 major species appeared after
- a second round of PCR amplification, the smallest of which was of the expected size. These fragments were cloned and sequenced. The smallest fragment did indeed have the sequence originally identified from the LifeSeq database, while the others contained
- insertions (figure 4).

As the first round of amplification suggested that the major species found in cDNA from ovary and placenta was not that originally identified in the LifeSeq<sup>TH</sup> database, the focus of effort was switched to the presumed major species (it seemed likely that

clones 57, 25-27 and 2.1kb clones 1-3 in fig 4 represented the major mRNA species). Conceptual. translation of the DNA sequences of these cloned PCR fragments indicated that the complete open reading 5 frame was not present in the clones or in the sequence from LifeSeq<sup>tM</sup>. While all clones contained the same sequence in the region of the translation termination codon, indicating that the end of the open reading frame had been identified, the 5' end of 10 the open reading frame had not been cloned. 5' RACE experiments were therefore carried out in order to find the start of the reading frame. PCR primers designed for RACE experiments are shown in figure 5. RACE PCR products were sequenced directly. Sequence 15 could be obtained from the 3' end of these RACE products but not from the 5' end; probably because the products were not cloned and were therefore heterogeneous at the 5' end. This new sequence was assembled with the existing cloned sequence to give 20 the sequence shown in figure 6. Searching the LifeSeq<sup>TM</sup> database with this sequence identifies ESTs which extend the sequence a further 140bp in the 5' direction and a further 160bp in the 3' direction (figure 7). This longer contig was used to design 25 oligonucleotide primers to amplify the entire coding sequence (these primer sequences are shown in figure PCR was carried out using primers 5'-1 and vegfX10 (in order to clone a "full-length" cDNA), and with primers 5'-1 and vegfX6 (in order to clone the 30 full coding region, see figure 3 for sequences of vegfX10 and vegfX6). A number of clones were obtained for the shorter fragment, of which clones 4 and 7 contain no PCR errors (sequence of clones 4 & 7 in figure 9). A single clone was obtained for the 35 longer fragment (clone 9), but this sequence appears to contain 2 PCR errors.

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The predicted polypeptide from these longer contigs is shown in figure 10. Amino acids 1-22 are predicted to encode a signal sequence (von Heijne, 5 1986, Nucleic Acids Res. 14, 4683-4690). Figure 11 shows an alignment of the protein sequence with VEGFs A-D. The region homologous to the other VEGFs is located towards the C-terminus of the protein. As the VEGF homology domain is expected to belong to the TGF-beta superfamily of growth factors and to consist 10 of a dimer containing both intra- and intermolecular disulphide bonds, initial alignments focussed on the cysteines. However, mapping of the sequence onto the known x-ray structure of the VEGF-A receptor-binding 15 domain (Muller et al (1997) Proc. Natl. Acad. Sci USA 94, 7192-7197) suggests that the alignment in figure 11 is plausible, as the extra 4 cysteine residues within the VEGF-homology region of VEGF-X (compared to this region of VEGF-A) correspond to residues which are spatially close in VEGF-A, and may 20 therefore be able to form disulphide bonds.

A search of the PFAM database of protein domains with the full-length polypeptide sequence from figure 10 25 identifies two domain consensus sequences within the polypeptide. The more C-terminal domain is a "VEGF" domain: (the known VEGFs all contain this domain and the structure of this region of VEGF-A is similar to that of PDGF). Additionally towards the N-terminus 30 of the polypeptide there is a CUB domain (amino acids -40-150). The CUB domain is a 100-110 amino acid extracellular domain found in a number of developmentally-regulated proteins. When the fulllength protein is used to search the protein 35 databases using the BLAST 2 algorithm, the scores for matches to CUB domain-containing proteins are more

significant than those to the other VEGFs. Interestingly, the most significant matches are to the CUB domains of Neuropilins, and Neuropilin-1 was recently identified as a receptor of one of the VEGF-A isoforms VEGF-A<sub>165</sub> (Soker et al. (1998) Cell 92, 735-745).

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Assuming that the variant sequences isolated by PCR (i.e. the smaller PCR fragments) use the same translation initiation site as the full-length 10 sequence, they would result in production of the variant proteins shown in figure 12. It may be significant that both of these variant proteins retain the CUB domain and delete all or part of the 15 VEGF-like domain. The production of these variant sequences can be explained by the use of a cryptic splice donor/acceptor site within the VEGF-X sequence (figure 30B, between nt. 998/999): one variant arises by splicing out of the region between nt. 729-998, the other by splicing out of the region between nt. 20 999-1187.

## Expression

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# 25 Full-length expression constructs Mammalian cells

Clone 4 containing the full CDS of VEGF-X (see figure 9), was used to generate constructs for expression of full-length protein. The sequence was amplified by PCR and cloned into the vector pCDNA6/V5-His so as to add a C-terminal V5 epitope tag and His, tag. The DNA and polypeptide sequence in this vector is shown in figure 19. Transient expression in COS cells followed by western blotting and detection via an anti-V5 mAb demonstrates the secretion of a protein of ~50K into the medium in transfected cells only

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(figure 22A). This construct can also be used to generate VEGF-X expressing stable CHO cell lines.

## Baculovirus/Insect-cell expression system

- For expression in the baculovirus/insect cell system the DNA encoding the predicted mature VEGF-X polypeptide sequence was fused to a sequence encoding a signal derived from melittin, a secreted insect protein. An N-terminal 6His tag was also added to
- facilitate purification. The insert was then cloned into the baculovirus expression vector pFASTBAC. The DNA and polypeptide sequence of this construct is shown in figure 20. Infection of Trichoplusia ni Hi5 cells with this recombinant baculovirus results in
- the secretion of a protein of approximately 45K into the medium (data not shown).

#### E.coli

The coding region of VEGF-X has been cloned in a

variety of ways for expression as a secreted protein
in E.coli. A particularly useful expression clone
carries an N-terminal fusion to the E.coli
maltose-binding protein (MBP- derived from the
expression vector pMAL-p2, New England Biolabs) and a

C-terminal fusion to a 6His tag. The DNA and

C-terminal fusion to a 6His tag. The DNA and polypeptide sequence of this vector is shown in figure 21. Sequential purification of cell fractions on Ni-NTA resin and amylose resin allows the isolation of the expressed protein (see figure 22B).

## Expression of fragments

## **VEGF**

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The VEGF domain of VEGF-X has been expressed in *E.coli*. Similar domains from VEGF-A (Christinger et al. (1996) *PROTEINS: Structure, Function and Genetics* 26, 353-357), and VEGF-D (Achen et al (1998) *Proc.* 

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Natl. Acad. Sci USA 95, 548-553) have been shown to be capable of binding to the respective receptors. Expression of these domains was carried out using the bacterium E.coli. Additionally, the full-length protein: was expressed using the baculovirus/insect cell expression system. The oligonucleotide primers which have been obtained for these experiments are shown in figure 13. The construct directed expression in the bacterial cytoplasm, and as expected the protein was produced in insoluble form in inclusion bodies (the DNA and polypeptide sequence used for PDGF domain expression is shown in figure 24). Inclusion bodies were washed, solubilized with urea and the protein purified under denaturing conditions, before refolding by dialysis to remove the urea. Soluble protein was obtained, but shows little evidence of the disulphide bond linked dimers seen with material derived from animal cells (figure 25, compare with figure 22A & B). It is not clear

## CUB

The CUB domain has been expressed as a soluble secreted protein in *E.coli* (figure 26). The protein was purified by binding to Ni-NTA resin (figure 27) and assayed for activity on HUVECs in an in-vitro proliferation assay.

therefore whether this protein is correctly folded.

## Properties of the VEGF-X protein

- The transient mammalian cell expression system described above has been used to generate full-length VEGF-X protein, as shown by antibody detection following Western blotting (see figure 22A).
- 35 <u>Disulphide bond linked dimers</u>
  The other members of the PDGF family of growth

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factors, the PDGFs and VEGFs, all exist as dimers in which two monomers constituting the dimer are linked by interchain disulphide bonds. The x-ray structures of PDGF-BB (Oefner et al, 1992), and VEGF-A (Muller et al, 1997) are known and indicate that at least these two members of the family contain two interchain disulphide bonds. Practically this means that in SDS-PAGE analysis of these growth factors the presence of interchain disulphide bonds is shown by a large decrease in mobility in the absence of reducing agent (ie. the nonreduced dimer migrates more slowly through the gel than the reduced monomer). This effect was also expected for VEGF-X, and has been demonstrated for the material obtained from transient mammalian cell expression (figure 22A). In the case of the full length material produced in E. coli only some 10% of the total VEGF-X protein appears to be present as disulphide bond-linked dimers (figure 22B). However, these results provide evidence that the mammalian cell-derived protein is correctly folded, and that a portion of the E.coli-derived protein is too.

# Glycosylation

25 There are 3 predicted potential N-linked glycosylation sites within the VEGF-X protein: at residues 25, 55 and 254 of the polypeptide sequence. The predicted molecular mass of the mature VEGF-X protein is 40kDa, but SDS-PAGE and western blotting 30 (detection via an introduced C-terminal epitope tagsee figure 19) of the full-length protein expressed in COS cells gives a band slightly larger than the expected size (45-50kDa) as well as one at 25kDa (figure 22A). This smaller band is presumed to be a 35 C-terminal proteolysis fragment derived from the full-length molecule (controls from uninfected cells do not show this band), probably corresponding to a

cleavage between the CUB and VEGF domains. EndoH treatment of the preparation gives a slight mobility change for the full-length protein (figure 23), but for the smaller VEGF domain fragment there is a clear change, indicating that the predicted glycosylation site within the VEGF domain at residue 254 is indeed glycosylated.

Activity of proteins in cell-based assays

Protein samples were tested for activity in cell proliferation, cell migration and in-vitro angiogenesis assays. Active samples can also be tested in the in vivo matrigel mouse model of angiogenesis.

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### Full-length VEGF-X protein

Conditioned medium derived from COS cells transiently expressing VEGF-X (see figure 22A) displayed no detectable activity in any of the assays. However, as VEGF-X protein could only be detected in this preparation by Western blotting, and not by Coomassie-staining of gels, it is clearly present at very low levels and this may be the reason for the observed lack of activity in the cell proliferation, migration or in vitro angiogenesis tests.

### VEGF domain

The VEGF domain protein described above has been tested in cell proliferation (on a range of cell types), cell migration and in vitro angiogenesis assays and has failed to show activity in any of these tests. As suggested above, this may be due to incorrect folding of this protein.

# 35 <u>CUB domain</u>

The CUB domain protein at the highest dose tested

(lµg/ml) appears to inhibit proliferation of HUVECs in the absence of other stimulation (figure 28A & B). This effect is also seen following stimulation with the lowest VEGF-A<sub>165</sub> dose tested (lng/ml- figure 28C). The CUB domain of VEGF-X therefore appears to show antiproliferative activity on HUVECs, even in the presence of low VEGF-A<sub>165</sub> doses.

### Tissue distribution of mRNA

10 VEGF-A mRNA expression has been shown to be upregulated in a wide variety of human tumors (lung, breast, ovarian, colon, stomach, liver, pancreas, kidney, bladder and prostate- Takahashi et al, 1995). Tumor VEGF-A expression has been shown to correlate 15 with tumor growth rate, microvascular density and tumor metastasis (Takahashi et al, 1995). thus of interest to examine the mRNA expression patterns of VEGF-X. Accordingly, Northern blot analysis of mRNA derived from different tissues has 20 been carried out. The results indicate that although the VEGF-X mRNA is expressed at low levels, it is present in a wide range of tissues. PCR amplification of cDNA from a range of tissue sources supports this idea (figure 29A). The major mRNA 25 species is approximately 3.1kb in size. There is no significant upregulation seen in tumour cell lines or in tumour tissues tested (figure 29B), with the possible exception of the cell lines GI-117 (lung carcinoma) and SaOS-2 (osteosarcoma). The results of 30 these initial tissue distribution studies do not, therefore, provide evidence for upregulation of

### Genomic structure of the VEGF-X gene

A genomic BAC clone covering the 3' part of the VEGF-X locus was isolated by hybridisation screening

VEGF-X in tumour growth, as is seen with VEGF-A.

of nylon filters containing a human BAC library. Direct sequencing of this clone using oligonucleotide primers based on the VEGF-X cDNA sequence allowed the determination of several intron/exon boundaries (figure 30). Interestingly, the position of the mRNA splice site within the PDGF domain (nt 1187/1188 in figure 30B) is conserved with respect to those in the VEGF-A and VEGF-D genes (Tischer et al, 1991;

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### Materials & Methods

Rocchigiani et al, 1998).

PCR, Cloning, DNA sequence determination and BAC screening.

- All primers were purchased from Eurogentec, Seraing, Belgium. Insert-specific sequencing primers (15- and 16-mers) were designed by visual inspection of the DNA sequences. DNA was prepared on Qiagen-tip-20 columns or on Qiaquick spin columns (Qiagen GmbH, Düsseldorf,
- Germany) and recovered from the spin columns in 30µl Tris/EDTA-buffer (10mM TrisHCl pH 7.5, 1 mM EDTA (sodium salt)). Sequencing reactions were performed using BigDye<sup>TM</sup> Terminator Cycle Sequencing Ready Reaction kits (Perkin Elmer, ABI Division, Foster City, CA, USA) and
- were run on an Applied Biosystems 377 DNA sequencer (Perkin Elmer, ABI Division, Foster City, CA, USA).

  Polymerase chain reactions were carried out according to standard procedures (Ausubel et al, 1997). The PCR fragments were cloned into vectors pCR2.1
- (Invitrogen, Carlsbad, CA. USA) or pCR-TOPO
  (Invitrogen, NL) according to the manufacturer's instructions. One of those vectors, plasmid VEGFX/pCR2.1 1TOPO FL

was deposited on 1 March 1999 under Accession No.

35 LMBP 3925. After sequence determination, the inserts were cloned into the desired expression vectors (see

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figures 19, 20, 21, 24 & 26).

A human genomic BAC library (Genome Systems, Inc., St Louis, MI, USA) was screened by hybridisation to 5 oligonucleotides derived from the VEGF-X cDNA sequence, according to the manufacturer's instructions. BAC DNA was prepared using a Qiagen plasmid midi kit (Qiagen GmbH, Düsseldorf, Germany ) according to the manufacturer's instructions with 10 some modifications (after clearing of the lysate from chromosomal DNA, supernatants from individual preparations were pooled on a single column (tip 100), and after the 70 % EtOH wash, the pellet was resuspended overnight at 4°C in 100 µl TE). 20-mer 15 sequencing primers were designed based on the known cDNA sequence, and sequencing carried out as above.

# 5' RACE

In order to extend the cDNA clone in a 5' direction RACE reactions were carried out. Since it was known that the mRNA is present in placenta and skeletal muscle, Marathon-Ready<sup>TM</sup> placenta and skeletal muscle cDNAs were purchased from Clontech (Palo Alto CA.

USA) and used according to the manufacturer's instructions. DNA fragments were excised from agarose gels, purified using QiaQuick PCR purification columns (Qiagen GmbH, Düsseldorf, Germany) and sequenced directly.

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VEGF-X protein expression and purification DNA fragments encoding the desired protein sequences were amplified by PCR and cloned into appropriate expression vector systems.

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For mammalian cell expression, the full coding

sequence was cloned into the vector pcDNA6/V5-his (Invitrogen Leek, NL, see figure 19 for construct sequence), so as to add a C-terminal peptide tag to assist in detection and purification.

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For insect cell expression the sequence of the predicted mature polypeptide was initially amplified to add an N-terminal 6His peptide and then cloned into the pMelBacB vector (Invitrogen, Leek, NL) to add an insect cell signal sequence. The entire insert was then PCR-cloned into the vector pFASTBAC-1 (LifeTechnologies, Gaithersburg, MA, USA) for construction of a baculovirus according to the manufacturer's instructions.

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For E.coli expression, the coding region was PCR amplified to add a C-terminal 6His tag and then cloned into the vector pMAL-p2 (New England Biolabs, Beverly, MA, USA). The coding sequence of this construct is shown in figure 21). The protein was purified first on Ni-NTA resin (Qiagen GmbH, Düsseldorf, Germany) and then on amylose resin (New England Biolabs, Beverly, MA, USA), according to the manufacturers' instructions.

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DNA sequences encoding the CUB and VEGF domain fragments of VEGF-X were PCR amplified and cloned into pET22b and pET21a (Novagen, Madison, WI, USA) respectively. The CUB domain protein was prepared either from the periplasm or medium of induced cultures by standard methods (Ausubel et al, 1997). The protein was initially purified by precipitation with 20% ammonium sulphate. After overnight dialysis vs 20mM Tris Hcl pH7.5, 100mM NaCl to remove ammonium sulphate, the protein was further purified on Ni-NTA resin as described above. The VEGF domain protein was expressed in insoluble form, and preparation of

inclusion bodies was carried out using standard procedures (Ausubel et al 1997). Inclusion bodies were dissolved in 6M guanidine hydrochloride, 20mM Tris Hcl pH8.0, 200mM NaCl, 1mM 2-mercaptoethanol, and purified on Ni-NTA resin (Qiagen GmbH, Düsseldorf, Germany) according to the manufacturer's instructions. The protein was refolded by dialysis against several changes of buffer containing decreasing concentrations of denaturant.

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Analysis of protein glycosylation was carried out using EndoH (Roche Molecular Biochemicals, Brussels, BE) according to the manufacturer's instructions.

# 15 Cell Proliferation Assay

Human umbilical vein endothelial cells (HUVECs) (Clonetics, San Diego, CA.) were trypsinized with 0.05% trypsin/0.53mM EDTA (Gibco, Gaithersburg, MD.), resuspended in the EGM-2(Clonetics, San Diego, CA.), counted, and distributed in a 96-well tissue culture plate at 5,000 cells/well. Following cell attachment and monolayer formation (16 hours), cells were stimulated with various concentrations of truncated VEGF-X (CUB domain or VEGF domain) or dilutions of culture supernatants of the full-length VEGF-X (COS 7 or HEK293) in DMEM (Gibco, Gaithersburg, MD.) containing 0.5% to 2% FBS (HyClone, Logan, UT) as indicated. For human fetal dermal fibroblasts (American Type Culture Collection, Rockville, MD.), the growth medium was replaced by DMEM containing 0.1% BSA (Sigma, St. Louise, MO.) with or without various concentrations of truncated VEGF-X proteins. For HCASMC (Clonetics, San Diego, CA.), the medium was replaced by DMEM containing 0.5% FBS. The cells were treated for a further 24 hr-72 hr. For the measurement of proliferation, the culture media were

replaced with 100 µl of DMEM containing 5% FBS and 3

μCi/ml of [3H]-thymidine (Amersham, Arlington
Heights, IL.). Following pulse labeling, cells were
fixed with methanol/acetic acid (3:1, vol/vol) for 1
hour at room temperature. The cells were washed

5 twice with 250 μl/well of 80% methanol. The cells
were solubilized in 0.05% trypsin (100μl/well) for
30 minutes then in 0.5% SDS (100 μl/well) for another
30 minutes. Aliquots of cell lysates (180 μl) were
combined with 2 ml of scintillation cocktail (Fisher,
10 Springfiled, NJ) and the radioactivity of cell
lysates was measured using a liquid scintillation
counter (Wallac 1409). In each case, samples were
performed in quadruplicate.

# 15 Chemotaxis Assay

The chemotactic response of HUVECs was assayed using a 48-well modified Boyden chamber (NeuroProbe, Cabin John, MD.) and collagen-coated (0.1mg/ml type I collagen, Collaboratic Biomedical, Bedford, MA.) polycarbonate membrane filters with a pore diameter of 8 µm (NeuroProbe, Cabin John, MD.). Cell suspensions (15,000/well) were loaded to the upper

part of the chemotaxis chamber and stimulated for 4

hours with rhVEGF<sub>165</sub> (0.1-10 ng/ml) (Calbiochem, San Diego, CA.) or various concentrations of truncated VEGF-X (PDGF domain). Cells remaining on the top of the membrane were removed. Migration was assessed by counting the number of cells that migrated to the lower side of the filter membrane. The membrane was

fixed with 10% formaldehyde for 15 min, followed by staining with Gill's hemotoxylin III (Poly Scientific, Bay Shore, NY.). The assay was performed in triplicates and six independent high power fields per well were counted using a light microscope at 250

magnification. The results were expressed as the fold of unstimulated cells (EGM containing 0.1% BSA).

### In Vitro Angiogenesis Assay

In vitro angiogenesis in fibrin gels was quantitated using spheroids of human umbilical vein endothelial cells (Korff et al., 1998). To generate endothelial cell spheroids of defined size and cell number, a 5 specific number of cells (~ 800 cells per spheroid) was suspended in EGM-2 culture medium containing 20% methylcellulose (Sigma, St. Louis, MO.), seeded into nonadherent round-bottom 96-well plates. All 10 suspended cells in one well contributed to the formation of a single endothelial cell spheroid within 24 hours. A fibrin gel stock solution was prepared freshly prior to use by mixing 3mg/ml fibrinogen (Calbiochem, San Diego, CA.) in Medium 15 199(Gibco, Gaithersburg, MD.). Assays were performed in 24-well culture plates. The lml fibrinogen stock was mixed with 50 HUVEC spheroids and the corresponding test substance including rh-VEGF165 or various concentration of VEGF-X. 20 spheriod-containing fibrinogen was rapidly transferred into 24-well plates. Fifteen microliters of thrombin (100 NIH U/ml stock, Sigma, St. Louis, MO.) was added to the gel for the fibrin gel formation. The gel formation usually occurred within 25 30 seconds. After gel formation, lml/well of Medium 199 supplemented with 20% FBS, lmg/ml ε-aminocaproic acid (Calbiochem, San Diego, CA.) and antibiotics were added. The gel was incubated at 37°C (5%CO2, 95% air, 100% humidity). After 3 days, in vitro 30 angiogenesis was quantitated by measuring the length of the three longest capillary sprouts that had grown out of each spheroid (100% magnification), analyzing at least 10 spheroids per experimental group and

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Matrigel Mouse Assay

experiment.

The matrigel mouse assay is carried out as described by Passanti et al (1992).

Analysis of VEGF-X gene expression by RT-PCR analysis.

- Oligonucleotide primers VEGF-E2 and VEGF-X14 (figure 16; figure 5) were used for the specific PCR amplification of a 350 bp fragment from VEGF-X. PCR amplifications were performed on human multiple
- tissue cDNA (MTCTM) panels (Clontech human MTC panels I and II and human Tumor MTC panel) normalised to the mRNA expression levels of six different housekeeping genes. In addition, cDNA was made from different tumor cell cultures (Caco-2 colorectal
- adenocarcinoma; T-84 colorectal carcinoma; MCF-7 breast adenocarcinoma; T-47D breast ductal gland carcinoma; HT1080 bone fibrosarcoma; SaOS-2 osteosarcoma; SK-N-MC neuroblastoma; HepG2 hepatoblastoma; JURKAT T-cell leukemia and THP-1
- myelomonocytic leukemia). For the preparation of tumor cell cDNA, cells were homogenised and total RNA prepared using the RNeasy Mini kit (Qiagen GmbH, Hilden, Germany) according to manufacturer's instructions. 1 µg of total RNA was reverse
- transcribed using oligo(dT)15 as a primer and 50 U of Expand<sup>TM</sup> Reverse Transcriptase (Boehringer Mannheim, Mannheim, Germany) according to the manufacturer's instructions. FCR reactions with VEGF-X-specific or glyceraldehyde-3-phosphate dehydrogenase
- 30 (G3PDH)-specific primers were then performed on 1 μl of this cDNA. For all cDNAs, PCR reactions with VEGF-X specific primers were performed in a total volume of 50 μl, containing 5 μl (± 1 ng) of cDNA, 1x Advantage KlenTaq PCR reaction buffer, 0.2 mM dNTP,
- 250 nM of primers VEGF-E2 and VEGF-X14 and 1 µl of Advantage KlenTaq polymerase mix. Samples were heated

to 95°C for 30 s and cycling was done for 30 s at 95°C and 30 s at 68°C for 25, 30 or 35 cycles. Control reactions using specific primers that amplify a 1 kb fragment of the housekeeping gene G3PDH were also performed according to the manufacturer's instructions.

Northern blot analysis of VEGF-X.

Northern blots containing 2 µg of poly(A)-rich RNA 10 derived from different human tissues (Clontech Laboratories; MTNTM blot, MTNTM blot II and Cancer Cell Line  $MTN^{TM}$  blot) were hybridized according to the manufacturers instructions with a  $\alpha-[^{32}P]-dCTP$ random-priming labelled (Multiprime labelling kit, Roche Diagnostics) 293 bp specific VEGF-X fragment 15 (PinAI-StuI fragment including 92 bp of the 3' end coding region and 201 bp of the 3' untranslated region of VEGF-X). The blots were hybridized overnight at 68°C and final washes at high stringency 20 were at 68°C in 0.1x SSC/0.1 % SDS. The membranes were autoradiographed for 1 to 3 days with intensifying screens.

# Full length VEGF-X

The effect of full length VEGF-X on proliferation of HuVEC cells was determined by the <sup>3</sup>H-Thymidine incorporation assay. HuVEC cells were serum starved for 24 hours prior to treatment with the full length VEGF-X at the concentration range from 100 pg/ml-10 µg/ml. There was no effect of VEGF-X at 100 pg/ml-10 ng/ml on endothelial cell proliferation. At the higher concentrations of FL-VEGF-X (100 ng/ml and 1 µg/ml) there was a marked inhibition of endothelial cell proliferation. This is probably due to the very high endotoxin level in the samples. The VEGF-X sample was purified in order to decrease the

endotoxin level and is currently tested in the cell proliferation assay.

The Summary from Testing the CUB Domain 5 The effect of CUB domain on inhibition of HuVEC prolieration either serum- (2%), rh-VEGF or bFGFstimulated, was assessed by the 3H-Thymidine incorporation assay. Cells were serum starved followed by the treatment with the CUB domain and 10 various growth factors. Results showed that the CUB domain inhibited endothelial cell proliferation, either serum- (2%), rh-VEGF or bFGF-stimulated in a dose dependent manner with maximal inhibition at 10 μg/ml. There was approximately a 2-fold inhibition 15 of proliferation (at 10 µg/ml) of cells stimulated with VEGF and bFGF and nearly a 5-fold inhibition of cells stimulated with serum (2%). Results with the LDH assay showed that there was no cytotoxicity associated with the inhibition of cell proliferation 20 by the CUB domain.

Therefore, the N-terminus of the polypeptide from Figure 10 has been shown to possess a CUB domain. When database searches are carried out using the full-length coding sequence the best matches (i.e. for a BLAST search, those with the lowest probability score) are found with the CUB domain rather than with the VEGF-like domain. The best match from searching release 37 of the SWISSPROT database (Feb 1999) is to the CUB domain of a neuropilin from Xenopus laevis, and the matches to the CUB domains of human neuropilins 1 and 2 are also more significant than matches to the VEGFs.

This similarity is provocative, given the identification of neuropilin-1 and -2 as cellular receptors for the VEGF-A 165 (Stoker et al. 1998,

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reviewed in Neufeld et al. 1999). It is plausible therefore that VEGF-X could exert dual regulatory effects: via interaction with the tyrosine kinase VEGF-receptors mediated by the VEGF-like domain, as well as via interaction with VEGF isoforms or with the neurophilin receptors, mediated by the CUB domain.

To the best of our understanding the latter would be entirely novel, and searches on the most recent release of the Incyte database do not reveal any other proteins containing both CUB and VEGF-like domains. This arrangement of domains suggests possible positive or negative models of regulation:

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Positive- the VEGF-like domain is able to interact productively with the tyrosine kinase VEGF receptors giving activation, and the CUB domain is able to interact productively with the neuropilin receptor

20 giving activation.

Negative- the VEGF-like domain does not interact productively with the tyrosine kinase VEGF receptors, either preventing receptor dimerisation or blocking the VEGF binding sites. Further, the CUB domain does not interact productively with the neuropilin receptors, either preventing receptor activation or blocking the VEGF binding sites, or indeed by binding to VEGF isoforms and preventing their interaction with receptors.

TABLE 1

	ORIGINAL RESIDUE	EXEMPLARY SUBSTITUTIONS
	ALA	SER, THR
5	ARG	LYS
	ASN	HIS, SER
	ASP	GLU, ASN
	CYS	SER
•	GTN .	ASN, HIS
10	Grn	ASP, GLU
	GLY	ALA, SER
	HIS .	ASN, GLN
	ILE	LEU, VAL, THR
	LEU	ILE, VAL
15	LYS	ARG, GLN, GLU, THR
	MET	LEU, ILE, VAL
	PHE	LEU, TYR
	SER	THR, ALA, ASN
20	THR	SER, ALA
	TRP	ARG, SER
	TYR	PHE
	VAL	ILE, LEU ALA
	PRO	ALA

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5	Sequence ID No 1	corresponds to the amino acid sequence from position 23 to 345 of the amino acid sequence illustrated in Figure 10.
	Sequence ID No 2	is the amino acid sequence illustrated in Figure 10.
10	Sequence ID No 3	corresponds to the sequence from position 257 to 1291 of the nucleotide sequence illustrated in Figure 9.
15	Sequence ID No 4	corresponds to the polynucleotide sequence of VEGFX1 illustrated in Figure 3.
20	Sequence ID No 5	corresponds to the polynucleotide sequence of VEGFX2 illustrated in Figure 3.
25	Sequence ID No 6	corresponds to the polynucleotide sequence of VEGFX3 illustrated in Figure 3.
30	Sequence ID No 7	corresponds to the polynucleotide sequence of VEGFX4 illustrated in Figure 3.
25	Sequence ID No 8	corresponds to the polynucleotide sequence of VEGFX5 illustrated in Figure 3.
35	Sequence ID No 9	corresponds to the polynucleotide sequence of VEGFX6 illustrated in

Figure 3.

5	Sequence	ID No 10	corresponds to the polynucleotide sequence of VEGFX7 illustrated in Figure 3.
10	Sequence	ID No 11	corresponds to the polynucleotide sequence of VEGFX8 illustrated in Figure 3.
10	Sequence	ID No 12	corresponds to the polynucleotide sequence of VEGFX9 illustrated in Figure 3.
15	Sequence	ID No 13	corresponds to the polynucleotide sequence of VEGFX10 illustrated in Figure 3.
20	Sequence	ID No 14	corresponds to the polynucleotide sequence of VEGFX11 illustrated in Figure 4.
25	Sequence	ID No 15	corresponds to the polynucleotide sequence of VEGFX12 illustrated in Figure 4.
	Sequence	ID No 16	corresponds to the polynucleotide sequence of VEGFX13 illustrated in Figure 4.
30	Sequence	ID No 17	corresponds to the polynucleotide sequence of VEGFX14 illustrated in Figure 4.
35	Sequence	ID No 18	corresponds to the polynucleotide sequence 5'-1 in Figure 8.

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	Sequence ID No 19	corresponds to the polynucleotide sequence 5'-2 in Figure 8.
5	Sequence ID No 20	corresponds to the polynucleotide sequence of VEGFX6 illustrated in Figure 13.
10	Sequence ID No 21	corresponds to the polynucleotide sequence of VEGFX7 illustrated in Figure 13.
15	Sequence ID No 22	corresponds to the polynucleotide sequence of VEGFX8 illustrated in Figure 13.
15	Sequence ID No 23	corresponds to the polynucleotide sequence of VEGFX9 illustrated in Figure 13.
20	Sequence ID No 24	corresponds to the polynucleotide sequence of VEGBAC1 illustrated in Figure 13.
25	Sequence ID No 25	corresponds to the polynucleotide sequence of VEGBAC2 illustrated in Figure 13.
30	Sequence ID No 26	corresponds to a polypeptide having the amino acid sequence from amino acid position 40 to 150 of the sequence of Figure 10.
35	Sequence ID No 27	corresponds to a polypeptide having the amino acid sequence illustrated in Figure 26.
	Sequence ID No 28	corresponds to the sequence from

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> position 5 to 508 of the nucleotide sequence illustrated in Figure 26. corresponds to the nucleotide sequence from position 5 to 508 of the nucleotide sequence illustrated in Figure 26.

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Sequence ID No 29

Sequence ID No 30 corresponds to the sequence from position 214 to 345 of the nucleotide sequence illustrated in Figure 10.

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# **CLAIMS**

- A nucleic acid molecule encoding a VEGF-X protein or a functional equivalent, derivative or bioprecursor thereof, said protein comprising any of the sequences from position 23 to 345 of the amino acid sequence illustrated in Figure 10, or the complete sequence as illustrated in Figure 10.
- A nucleic acid molecule according to claim 1 wherein said nucleic acid is a DNA molecule.
  - 3. A nucleic acid molecule according to claim 1 or 2 wherein said nucleic acid is a cDNA molecule.
- A nucleic acid molecule according to claim 3 comprising the nucleotide sequence from position 257 to 1291 of the nucleotide sequence illustrated in Figure 9, or sequences that hybridise thereto under high stringency conditions or the complement thereto.
  - 5. An antisense molecule capable of hybridising to a molecule according to any of claims 1 to 4 under high stringency conditions.
  - 6. A nucleic acid molecule according to any of claims 1 to 4 which is of mammalian origin.
- 7. A nucleic acid molecule according to claim 6 which is of human origin.
  - 8. An isolated VEGF-X protein, or a functional equivalent, derivative or bioprecursor thereof, having an amino acid sequence from position 23 to 345 of the amino acid sequence illustrated in Figure 10 or the complete amino acid sequence of Figure 10.

9. A VEGF-X protein, or a functional equivalent, derivative or bioprecusor thereof, encoded by a nucleic acid molecule as defined in any of claims 1 to 4.

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- 10. A protein according to claim 9, which comprises the amino acid sequence illustrated in Figure 10.
- 11. An expression vector comprising a nucleic acid molecule according to any of claims 1 to 4.
  - 12. An expression vector according to claim 11 further comprising a nucleotide sequence encoding a reporter molecule.

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- 13. An expression vector comprising an antisense molecule according to claim 5.
- 14. A nucleic acid molecule according to any of claims 1 to 4 or an antisense molecule according to claim 5 for use as a medicament.
  - 15. A host cell transformed or transfected with an expression vector according to claim 11 or 12.

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- 16. A host cell transformed or transfected with an expression vector according to claim 13.
- 17. A transgenic cell, tissue or organism comprising a transgene capable of expressing a VEGF-X protein according to claim 8 or 9.
  - 18. A transgenic cell, tissue or organism according to claim 17, wherein said transgene is included in an expression vector.
    - 19. A VEGF-X protein or a functional equivalent,

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derivative or bioprecursor thereof, expressed by a cell according to claim 15.

- 20. A VEGF-X protein, or a functional equivalent, derivative or bioprecursor thereof, expressed by a transgenic cell, tissue or organism according to claim 17.
- 21. A process for producing a VEGF-X protein
  according to any of claims 8 to 10, said process
  comprising transforming a host cell or organism with
  an expression vector according to claim 11, and
  recovering the expressed protein from said host cell
  or organism.
- 22. An antibody capable of binding to a protein according to any of claims 8 to 10, or an epitope thereof.

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- 20 23. An antibody according to claim 22 for use as a medicament.
- 24. A pharmaceutical composition comprising an antibody according to claim 22 together with a pharmaceutically acceptable carrier diluent or excipient thereof.
- 25. A method of identifying VEGF-X protein in a sample which method comprises contacting said sample with an antibody according to claim 22 and monitoring for binding of any protein to said antibody.
- 26. A kit for identifying the presence of VEGF-X protein in a sample which comprises an antibody
  35 according to claim 22 and means for contacting said antibody with said sample.

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- 27. A method of identifying compounds which modulate angiogenesis which method comprises providing a host cell or organism according to claim 15 or a transgenic cell, tissue or organism according to claim 17, contacting a test compound with said cell, tissue or organism and monitoring for an effect of said compound on said VEGF compared to a host cell or organism according to claim 15 or a transgenic cell
- 10 not been contacted with said compound.

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28. A compound identifiable according to the method of claim 27.

tissue or organism according to claim 17 which has

- 29. A compound according to claim 28 for use as a medicament.
- 30. A nucleic acid sequence comprising the nucleotide sequences illustrated in any of Figures 3,5, 8 or 13.
  - 31. A method for producing a polypeptide, said method comprising the steps of:
- 25 a) culturing the host cell of claim 15 under conditions suitable for expression of the polypeptide; and
  - b) recovering the polypeptide from the host cell culture.

32. A method of inhibiting angiogenic activity and inappropriate vascularisation including formation and proliferation of new blood vessels, growth and development of tissues, tissue regeneration and organ and tissue repair in a subject said method comprising administering to said subject an amount of an antisense molecule according to claim 5 in sufficient

concentration to reduce or prevent said angiogenic activity.

- 33. A method of inhibiting angiogenic activity or inappropriate vascularisation including any of formation and proliferation of new blood vessels, growth and development of tissues, tissue regeneration and organ and tissue repair in a subject said method comprising administering to said subject an amount of an antibody according to claim 22 in sufficient concentration to reduce or prevent said angiogenic activity or inappropriate vascularisation.
- 34. A method of inhibiting angiogenic activity or inappropriate vascularisation including any of formation and proliferation of new blood vessels, growth and development of tissues, tissue regeneration and organ and tissue repair in a subject, said method comprising implanting in said subject cells that express an antibody according to claim 22.
- 35. A method of treating or preventing any of cancer, rheumatoid arthritis, psoriasis and diabetic retinopathy, said method comprising administering to said subject an amount of an antisense molecule according to claim 5 in sufficient concentration to treat or prevent said disorders.
- 36. A method of treating or preventing any of cancer, rheumatoid arthritis, psoriasis and diabetic retinopathy, said method comprising administering to said subject an amount of an antibody according to claim 22 in sufficient concentration to reduce or prevent said disorders.
  - 37. A method of promoting angiogenic activity or

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vascularisation to promote wound healing, skin graft growth, tissue repair, proliferation of new blood vessels, tissue regeneration and organ repair which method comprises applying or delivering to a site of interest a therapeutically effective amount of any of a group selected from a protein according to claim 8 and a nucleic acid molecule encoding a VEGF-X protein or a functional equivalent, derivative or bioprecursor thereof comprising an amino acid sequence illustrated in Figure 10, an expression vector comprising said nucleic acid molecule and a pharmaceutical composition comprising any of said nucleic acid molecule and said protein.

- 38. A method of treating wounds selected from the group consisting of dermal ulcers, pressure sores, venous sores, diabetic ulcers and burns by applying to said wound a therapeutically effective amount of any of a VEGF-X protein according to claim 8, a pharmaceutical composition comprising said protein and a pharmaceutically acceptable carrier, diluent or excipient therefor.
- 39. A nucleic acid molecule encoding a polypeptide having a CUB domain said polypeptide comprising the amino acid sequence from position 40 to 150 of the sequence of Figure 10.
- 40. A nucleic acid molecule encoding a polypeptide 30 having a CUB domain, said polypeptide comprising the amino acid sequence of Figure 26.
- 41. A nucleic acid molecule according to claim 39 or
  40, comprising the nucleotide sequence from position
  5 to 508 of the sequence illustrated in Figure 26.
  - 42. A nucleic acid molecule according to any of

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claims 39 to 41 comprising the nucleotide sequence illustrated in Figure 26.

- 43. A nucleic acid molecule encoding a VEGF like
  5 domain comprising the sequence from position 214-345
  of the sequence of Figure 10 or the sequence from
  position 15 to 461 illustrated in Figure 24.
- 44. An expression vector comprising a nucleic acid molecule according to any of claims 39 to 42.
  - 45. An expression vector comprising a nucleic acid molecule according to claim 43.
- 46. A host cell transformed or transfected with an expression vector according to claim 44.
  - 47. A host cell transformed or transfected with an expression vector according to claim 45.
  - 48. A protein expressed by the cell according to claim 46.
- 49. A protein expressed by the cell according to claim 47.
  - 50. A method of identifying compounds that inhibit or enhance angiogenic activity, said method comprising contacting a cell expressing a VEGF
- receptor and/or a neuropilin 1 or 2 type receptor with said compound in the presence of a VEGF-X protein according to claim 8 and monitoring for the effect of said compound or said cell when compared to a cell which has not been contacted with said compound.
  - 51. A compound identifiable according to the method

of claim 50 as an inhibitor or enhancer of angiogenic activity.

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- 52. A method of inhibiting angiogenic activity or 5 inappropriate vascularisation, said method comprising contacting a cell expressing a VEGF receptor and a neuropilin type receptor with a protein selected from any of a protein according to any of claims 8 to 10 and a protein according to claim 48 or a protein 10 according to claim 49.
  - 53. Use of a nucleotide sequence illustrated in any of Figures 14 and 15 in identifying a VEGF-X protein according to claim 8.
- 15 A nucleic acid molecule encoding a polypeptide comprising a CUB domain having the sequence from position 40 to 150 of the sequence of Figure 10 or from position 5 to 508 of the sequence of Figure 26 20 and a sequence encoding a VEGF domain.
  - A nucleic acid molecule according to claim 54 wherein said sequence encoding said VEGF domain is selected from the sequences encoding any of VEGF A to D or isoforms or variants thereof.
  - A nucleic acid molecule encoding a polypeptide comprising the amino acid sequence from position 40 to 150 of the sequence illustrated in Figure 10 for use as a medicament.

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Use of a nucleic acid molecule encoding a polypeptide having the amino acid sequence from position 40 to 150 of the sequence illustrated in 35 Figure 10 in the manufacture of a medicament for treatment of disease conditions associated with inappropriate angiogenesis such as tumour or cancer WO 00/37641

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growth, retinopathy, osteoarthritis or psoriasis.

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58. A polypeptide comprising the amino acid sequence from position 40 to 150 of the sequence illustrated in figure 10 for use as a medicament.

- 59. A polypeptide comprising the amino acid sequence from position 40 to 150 of the sequence illustrated in Figure 10 in the manufacture of a medicament for the treatment of disease conditions associated with inappropriate angiogenesis such as tumour growth, retinopathy, osteoarthritis or psoriasis.
- 60. Use of a CUB domain comprising the amino acid sequence from position 40 to 150 of the sequence of Figure 10, or the amino acid sequence of Figure 26, to identify compounds which inhibit angiogenic activity in a method according to claim 50.
- 20 61. A method of inhibiting angiogenic activity and inappropriate vascularisation including formation and proliferation of new blood vessels, growth and development of tissues, tissue regeneration and organ and tissue repair in a subject said method comprising administering to said subject an amount of a polypeptide having an amino acid sequence from position 40 to 150 of the sequence illustrated in Figure 10 or a nucleic acid molecule according to any of claims 39 to 42 in sufficient concentration to reduce or prevent said angiogenic activity.
- 62. A method of treating or preventing any of cancer, rheumatoid arthritis, psoriasis and diabetic retinopathy, said method comprising administering to said subject an amount of a polypeptide having an amino acid sequence from position 40 to 150 of the sequence illustrated in Figure 10 or a nucleic acid

molecule according to any of claims 39 to 42 in sufficient concentration to treat or prevent said disorders.

- 5 63. An antisense molecule capable of hybridising to a molecule according to any of claims 39 to 42 under high stringency conditions.
- 64. An antisense molecule capable of hybridising to a molecule according to claim 43 under high stringency conditions.
- 65. A transgenic cell, tissue or organism comprising a transgene capable of expressing a protein according to claim 48.
  - 66. A transgenic cell, tissue or organism comprising a transgene capable of expressing a protein according to claim 49.

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67. A transgenic, cell tissue or organism according to claim 65 or 66, wherein said transgene is included in an expression vector according to claim 41 or 42.

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- 68. An antibody capable of binding to a protein according to claim 48 or an epitope thereof.
- 69. An antibody capable of binding to a protein according to claim 49 or an epitope thereof.
  - 70. A pharmaceutical composition comprising an antibody according to claim 68 or 69 together with a pharmaceutically acceptable carrier diluent or excipient therefor.
  - 71. A pharmaceutical composition comprising a

compound according to claim 48 together with a pharmaceutically acceptable carrier, diluent or excipient therefor.

72. A nucleic acid molecule encoding a variant of a VEGF-X protein having any of the sequences of nucleotides illustrated in Figure 12.

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FIGI

AAAATGTATG GATACAACTT ACGTTTGATG AAAGATTTGG GCTTGAAGAC CCAGAAGATG TTTTACATAC CTATGTTGAA TGCAAACTAC TTTCTAAACC CGAACTTCTG GGTCTTCTAC ACATATGCAA GTATGATTTT GTAGAAGTTG AGGAACCCAG TGATGGAACT ATATTAGGGC TGTATACGTT CATACTAAAA CATCTTCAAC TCCTTGGGTC ACTACCTTGA TATAATCCCG 121 GCTGGTGTGG TTCTGGTACT GTACCAGGAA AACAGATTTC TAAAGGAAAT CAAATTAGGA CGACCACAC AAGACCATGA CATGGTCCTT TTGTCTAAAG ATTTCCTTTA GTTTAATCCT +1 MetAsn IlePheLeu LeuAsnLeuLeu ThrGluGlu ValArgLeu ]-----TAAGATTTGT ATCTGATGAA TATTTTCCTT CTGAACCTTC TAACAGAGGA GGTAAGATTA ATTCTAAACA TAGACTACTT ATAAAAGGAA GACTTGGAAG ATTGTCTCCT CCATTCTAAT +1 TyrSerCysThr ProArgAsn PheSerVal SerIleArgGlu GluLeuLys ArgThrAsp TACAGCTGCA CACCTCGTAA CTTCTCAGTG TCCATAAGGG AAGAACTAAA GAGAACCGAT ATGTCGACGT GTGGAGCATT GAAGAGTCAC AGGTATTCCC TTCTTGATTT CTCTTGGCTA +1 ThrIlePheTrp ProGlyCys LeuLeuVal LysArgCysGly GlyAsnCys AlaCysCys 301 ACCATTTCT GGCCAGGTTG TCTCCTGGTT AAACGCTGTG GTGGGAACTG TGCCTGTTGT TGGTAAAAGA CCGGTCCAAC AGAGGACCAA TTTGCGACAC CACCCTTGAC ACGGACAACA +1 LeuHisAsnCys AsnGluCys GlnCysVal ProSerLysVal ThrLysLys TyrHisGlu -----361 CTCCACAATT GCAATGAATG TCAATGTGTC CCAAGCAAAG TTACTAAAAA ATACCACGAG GAGGTGTTAA CGTTACTTAC AGTTACACAG GGTTCGTTTC AATGATTTTT TATGGTGCTC +1 ValLeuGlnLeu ArgProLys ThrGlyVal ArgGlyLeuHis LysSerLeu ThrAspVal GTCCTTCAGT TGAGACCAAA GACCGGTGTC AGGGGATTGC ACAAATCACT CACCGACGTG CAGGAAGTCA ACTCTGGTTT CTGGCCACAG TCCCCTAACG TGTTTAGTGA GTGGCTGCAC +1 AlaLeuGluHis HisGluGlu CysAspCys ValCysArgGly SerThrGly Gly ------GCCCTGGAGC ACCATGAGGA GTGTGACTGT GTGTGCAGAG GGAGCACAGG AGGATAGCCG CGGGACCTCG TGGTACTCCT CACACTGACA CACACGTCTC CCTCGTGTCC TCCTATCGGC 601 ACGTATGCGT TATCTCCATC CTTAATCTCA GTTGTTTGCT TCAAGGACCT TTCATCTTCA TGCATACGCA ATAGAGGTAG GAATTAGAGT CAACAAACGA AGTTCCTGGA AAGTAGAAGT 661 GGATTTACAG TGCATTCTGA AAGAGGAGAC ATCAAACAGA ATTAGGAGTT GTGCAACAGC CCTAAATGTC ACGTAAGACT TTCTCCTCTG TAGTTTGTCT TAATCCTCAA CACGTTGTCG TCTTTTGAGA GGAGGCCTAA AGGACAGGAG AAAAGGTCTT CAATCGTGGA AAGAAAATTA AGAAAACTCT CCTCCGGATT TCCTGTCCTC TTTTCCAGAA GTTAGCACCT TTCTTTTAAT 781 AATGTTGTAT TAAATAGATC ACCAGCTAGT TTCAGAGTTA CCATGTACGT ATTCCACTAG TTACAACATA ATTTATCTAG TGGTCGATCA AAGTCTCAAT GGTACATGCA TAAGGTGATC

	FIG	1 (CONTINU	رحم <u>)</u>			
0.41						
841		TATTTCAGTT				
	GACCCAAGAC	ATAAAGTCAA	GAAAGCTATG	CCGAATCCCA	TTACAGTCAT	GTCCTTTTTT
901	ACTGTGCAAG	TGAGCACCTG	ATTCCGTTGC	CTTGCTTAAC	TCTAAAGCTC	CATGTCCTGG
		ACTCGTGGAC				
961	GCCTAAAATC	GTATAAAATC	TGGATTTTTT	THETTE	TGCTCATATT	CACATATGTA
		CATATTTTAG				
1021						
1021		TTCTATGTAC				
		AAGATACATG		•		
1081		GTCGTGCTGA				
	AATTTGAACA	CAGCACGACT	ATCCTGTCTG	ACCTAAAAAG	TATAAAGAAT	AATTTTAAAG
1141	TGCCATTTAG	AAGAAGAGAA	CTACATTCAT	GGTTTGGAAG	AGATAAACCT	GAAAAGAAGA
		TTCTTCTCTT				
1201		CTTCACTTTA				
		GAAGTGAAAT				
1261						
1201		TTGACATTAT				
		AACTGTAATA				
1321		ATTTAATATT				
		TAAATTATAA				•
1381	AAATTTTTCT	AAACACAATT	GTTATAGCCA	GAGGAACAAA	GATGATATAA	AATATTGTTG
	TTTAAAAAGA	TTTGTGTTAA	CAATATCGGT	CTCCTTGTTT	CTACTATATT	TTATAACAAC
1441	CTCTGACAAA	AATACATGTA	TTTCATTCTC	GTATGGTGCT	AGAGTTAGAT	TAATCTGCAT
	GAGACTGTTT	TTATGTACAT	AAAGTAAGAG	CATACCACGA	TCTCAATCTA	ATTAGACGTA
1501	TTTAAAAAAC	TGAATTGGAA	TAGAATTGGT	AAGTTGCAAA	GACTTTTTGA	AAATAATTAA
		ACTTAACCTT				
1561		CTTCCATTCC				
						ATACTTTCAT
1621	GACATTCAGA	TCCAGCCATT	ACTAACCTAT	TCCTTTTTTG	GGGAAATCTG	AGCCTAGCTC
						TCGGATCGAG
1681	AGAAAAACAT	AAAGCACCTT	GAAAAAGACT	TGGCAGCTTC	CTGATAAAGC	GTGCTGTGCT
						CACGACACGA
						01100110110
1741	GTGCAGTAGG	AACACATCCT	ATTTATTGTG	ATGTTGTGGT	тттаттатст	TAAACTCTGT
						ATTTGAGACA
1801						TCTTAACCAG
~ • • • • • • • • • • • • • • • • • • •						
	AGIAIGIGA	ACATATTAT	GTACCTATAA	AAATACATGT	CTTCATACAG	AGAATTGGTC
1861	TTCACTTATT	GTACCTGG				
	AAGTGAATAA					

vegfX8

vegfX9

vegfX10

# FIG. 2. Predicted VEGF-like protein encoded by Incyte contig of 8/12/98

- 1 MNIFLLNLLT EEVRLYSCTP RNFSVSIREE LKRTDTIFWP GCLLVKRCGG
- 51 NCACCLHNCN ECQCVPSKVT KKYHEVLQLR PKTGVRGLHK SLTDVALEHH
- 101 EECDCVCRGS TGG

# PCR primers for cloning VEGF-X vegfX1 AAAATGTATGGATACAACTTAC vegfX2 GTTTGATGAAAGATTTGGGCTTG vegfX3 TTTCTAAAGGAAATCAAATTAG vegfX4 GATAAGATTTGTATCTGATG vegfX5 GATGTCTCCTCTTTCAG vegfX6 GCACAACTCCTAATTCTG vegfX7 AGCACCTGATTCCGTTGC

TAGTACATAGAATGTTCTGG

AAGAGACATACTTCTGTAC

CCAGGTACAATAAGTGAACTG

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FIG. 4. Variants isolated by PCR (at 8/2/99, all cloned and sequenced at JRF)

a b

c d

**.** 

PCR primers- → →

\_ \_

← ←

Incyte contig \_\_\_

(8/12/98)

clone 22, 29, 41

9. 41

clone 52, 59

clone 15, 20

clones 57, 25,

26, 27

2.1kb clones 1,

2, 3

primers-

a- vegfX1

b- vegfX2

c- vegfX5

(see fig 3)

d- vegfX6

e- vegfX9

f-vegfX10

# FIG. 5. VEGF-X 5' RACE primers

vegfX11	CCTTTAGAAATCTGTTTTCCTGGTACAG
vegfX12	GGAAAATATTCATCAGATACAAATCTTATCC
vegfX13	GGTCCAGTGGCAAAGCTGAAGG
vegfX14	CTGGTTCAAGATATCGAATAAGGTCTTCC

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# FIG. 6. DNA sequence assembled from in-house clones and 5'RACE

1				TGCAGCCTTC		
	ACGGTCTCGT	CCACCGCGA	AGGTGGGGTC	ACGTCGGAAG	GGGACCGCCA	CCACTTTCTC
61				CCGTGAGTGA		
	TGAGCCCTCA	GCGACGAAGG	TTTCACGGGC	GGCACTCACT	CGAGAGTGGG	GTCAGTCGGT
+2		PheGlyLeuLe	eu LeuLeuTh	r SerAlaLeu	AlaGlyGlnA	rg GlnGlyTh
121	AATGAGCCTC	TTCGGGCTTC	TCCTGCTGAC	ATCTGCCCTG	GCCGGCCAGA	GACAGGGGAC
	TTACTCGGAG	AAGCCCGAAG	AGGACGACTG	TAGACGGGAC	CGGCCGGTCT	CTGTCCCCTG
+2	rGlnAlaGlu	SerAsnLeuSe	er SerLysPh	e GlnPheSer	SerAsnLysG	lu GlnAsnGl
181				CCAGTTTTCC		
ن .	AGTCCGCCTT	AGGTTGGACT	CATCATTTAA	GGTCAAAAGG	TCGTTGTTCC	TTGTCTTGCC
+2	yValGlnAsp		-	e ThrValSer	<del>-</del>	
241				TACTGTGTCT		
	TCATGTTCTA	GGAGTCGTAC	TCTCTTAATA	ATGACACAGA	TGATTACCTT	CATAAGTGTC
+2	rProArgPhe	ProHisThrT	yr ProArgAs	n ThrValLeu	ValTrpArgL	eu ValAlaVa
301	CCCAAGGTTT	CCTCATACTT	ATCCAAGAAA	TACGGTCTTG	GTATGGAGAT	TAGTAGCAGT
	GGGTTCCAAA	GGAGTATGAA	TAGGTTCTTT	ATGCCAGAAC	CATACCTCTA	ATCATCGTCA
+2	lGluGluAsn	ValTrpIleG	ln LeuThrPh	e AspGluArg	PheGlyLeuG	lu AspProGl
361	AGAGGAAAAT	GTATGGATAC	AACTTACGTT	TGATGAAAGA	TTTGGGCTTG	AAGACCCAGA
	TCTCCTTTTA	CATACCTATG	TTGAATGCAA	ACTACTTTCT	AAACCCGAAC	TTCTGGGTCT
+2	uAspAspIle	CysLysTyrA	sp PheValGl	u ValGluGlu	ProSerAspG	ly ThrIleLe
421				AGTTGAGGAA		
	TCTACTGTAT	ACGTTCATAC	TAAAACATCT	TCAACTCCTT	GGGTCACTAC	CTTGATATAA
+2	uGlyArgTrp	CysGlySerG	ly ThrValPr	o GlyLysGln	IleSerLysG	ly AsnGlnIl
481	AGGGCGCTGG	TGTGGTTCTG	GTACTGTACC	AGGAAAACAG	ATTTCTAAAG	GAAATCAAAT
	TCCCGCGACC	ACACCAAGAC	CATGACATGG	TCCTTTTGTC	TAAAGATTTC	CTTTAGTTT
+2	eArgIleArg	PheValSerA	sp GluTyrPh	e ProSerGlu	ProGlyPheC	ys IleHisTy
541				TCCTTCTGAA		
	ATCCTATTCT	AAACATAGAC	TACTTATAAA	AGGAAGACTT	GGTCCCAAGA	CGTAGGTGAT
+2	rAsnIleVal	MetProGlnP	he ThrGluAl	a ValSerPro	SerValLeuP	ro ProSerAl
601	CAACATTGTC	ATGCCACAAT	TCACAGAAGO	TGTGAGTCCT	TCAGTGCTAC	CCCCTTCAG
	GTTGTAACAG	TACGGTGTTA	AGTGTCTTCG	ACACTCAGGA	AGTCACGATG	GGGGAAGTC
+2	aLeuProLeu	AspLeuLeuA	sn AsnAlaIl	e ThrAlaPhe	SerThrLeuG	lu AspLeuI
661	TTTGCCACTG	GACCTGCTTA	ATAATGCTAT	AACTGCCTTT	AGTACCTTGG	AAGACCTTA'

	F/6 61	CONTINUE	חצו			
+2	eArgTyrLeu			ı AspLeuGlu	AspLeuTyrA	g ProThrTr
721		GAACCAGAGA CTTGGTCTCT				
+2	pGlnLeuLeu	GlyLysAlaPl	ne ValPheGly	y ArgLysSer	ArgValValAs	sp LeuAsnLe
781		GGCAAGGCTT CCGTTCCGAA				
+2	uLeuThrGlu	GluValArgLe	eu TyrSerCys	s ThrProArg	AsnPheSerVa	al SerIleAr
841		GAGGTAAGAT CTCCATTCTA				
	gGluGluLeu	•				
901		AAGAGAACCG TTCTCTTGGC				
+2	sGlyGlyAsn	CysAlaCysCy	/s LeuHisAsı	n CysAsnGlu	CysGlnCysVa	al ProSerLy
961	TGGTGGGAAC	TGTGCCTGTT	GTCTCCACAA	TTGCAATGAA	TGTCAATGTG	TCCCAAGCAA
		ACACGGACAA				
+2	sValThrLys	LysTyrHisG:	lu ValLeuGli	n LeuArgPro	LysThrGlyVa	al ArgGlyLe
1021		AAATACCACG TTTATGGTGC				
+2	uHisLysSer	LeuThrAspVa	al AlaLeuGl	ı HisHisGlu	GluCysAspCy	ys ValCysAr
1081		CTCACCGACG GAGTGGCTGC				
+2	gGlySerThr					
1141	AGGGAGCACA TCCCTCGTGT	GGAGGATAGC CCTCCTATCG	CGCATCACCA GCGTAGTGGT	CCAGCAGCTC GGTCGTCGAG	TTGCCCAGAG AACGGGTCTC	CTGTGCAGTG GACACGTCAC
1201	CAGTGGCTGA					
	GTCACCGACT	AAGATAATCT	CTTGCATACG	CAATAGAGGT	AGGAATTAGA	GTCAACAAAC
1261	CTTCAAGGAC GAAGTTCCTG	CTTTCATCTT GAAAGTAGAA				
1321	GAATTAGGAG					
	CTTAATCCTC	AACACGTTGT	CGAGAAAACT	CTCCTCCGGA	TTTCCTGTCC	TCTTTTCCAG
1381	TTCAATCGTG AAGTTAGCAC	GAAAGAAAAT CTTTCTTTTA				
1441	TACCATGTAC					
	ATGGTACATG	CATAAGGTGA	ŢCGACCCAAG	ACATAAAGTC	AAGAAAGCTA	TGCCGAATCC
1501	GTAATGTCAG	TACAGGAAAA	AAACTGTGCA	AGTGAGCACC	TGATTCCGTT	GCCTTGCTTA

	F16.61	CONTINUEL	0 <i>2</i> ).			
1561	ACTCTAAAGC			TCGTATAAAA	TCTGGATTTT	TTTTTTTTT
	TGAGATTTCG	AGGTACAGGA	CCCGGATTTT	AGCATATTTT	AGACCTAAAA	AAAAAAAAA
1621	TTTGCTCATA	TTCACATATG	TAAACCAGAA	CATTCTATGT	ACTACAAACC	TGGTTTTTAA
	AAACGAGTAT	AAGTGTATAC	ATTTGGTCTT	GTAAGATACA	TGATGTTTGG	ACCAAAAATT
1681	AAAGGAACTA	TGTTGCTATG	AATTAAACTT	GTGTCGTGCT	GATAGGACAG	ACTGGATTTT
	TTTCCTTGAT	ACAACGATAC	TTAATTTGAA	CACAGCACGA	CTATCCTGTC	TGACCTAAAA
1741					AACTACATTC	
					TTGATGTAAG	
1801	AGAGATAAAC					
					ATAGCTATTC	
1861					ATAACTGTTG	
1001					TATTGACAAC	
1921					TTCTTTTTA AAGAAAAAAT	
1981					TTGTTATAGC	
1901					AACAATATCG	
	CIAGIIGAIA	AAAATCGAAC	CATTIAAAAA	GATTIGIGIT	AACAATATCG	GICICCIIGI
2041	AAGATGATAT	AAAATATTGT	TGCTCTGACA	AAAATACATG	TATTTCATTC	TCGTATGGTG
					ATAAAGTAAG	
2101	CTAGAGTTAG	ATTAATCTGC	ATTTTAAAAA	ACTGAATTGG	AATAGAATTG	GTAAGTTGCA
	GATCTCAATC	TAATTAGACG	TAAAATTTTT	TGACTTAACC	ТТАТСТТААС	CATTCAACGT
2161					CCTGTTATTG	
	•				GGACAATAAC	
2221					TTACTAACCT	
2201					AATGATTGGA	
2281					TTGAAAAAGA	
2241						GAACCGTCGA
2341	TCCTGATAAA					
2401						ACTACAACAC
2401	GTTTTATTAT					
2461			CAAGGTATGT	GAACATATTT	ATGTACCTAT	AAAAATACAT
2401	CAGAAGTATG GTCTTCATAC			•		
	GICTICATAC	AUAUA				

# FIG. 7. New Sequence + Incyte ESTs

1		ACCTTGGGAA TGGAACCCTT				
61		CAGAAGAGGG GTCTTCTCCC				
121		TCTCTGCTGC				
101		AGAGACGACĢ TGAAAGAGAC		•		
181	_	ACTTTCTCTG				
+2					LeuLeuThrSe	
241		GTCAGCCAAA CAGTCGGTTT	TGAGCCTCTT	CGGGCTTCTC	CTGCTGACAT	CTGCCCTGGC
+2	aGlyGlnArg	GlnGlyThrGl	ln AlaGluSe	AsnLeuSer	SerLysPheG	ln PheSerSe
301	_	CAGGGGACTC GTCCCCTGAG				
+2	rAsnLysGlu	GlnTyrGlyVa	al GlnAspPro	GlnHisGlu	ArgIleIleTh	r ValSerTh
361		CAGTACGGAG GTCATGCCTC				
+2	rAsnGlySer	IleHisSerPr	o ArgPhePro	HisThrTyr	ProArgAsnTh	ır ValLeuVa
421	•	ATTCACAGCC TAAGTGTCGG				
+2	lTrpArgLeu	ValAlaValG	u GluAsnVal	TrpIleGln	LeuThrPheAs	sp GluArgPh
481		GTAGCAGTAG CATCGTCATC				
+2	eGlyLeuGlu	AspProGluAs	sp AsplleCys	LysTyrAsp	PheValGluVa	al GluGluPr
541		GACCCAGAAG CTGGGTCTTC				-
+2	oSerAspGly	ThrIleLeuGl	y ArgTrpCys	GlySerGly		y LysGlnIl
601		ACTATATTAG TGATATAATC				
+2	eSerLysGly	AsnGlnIleAr	g IleArgPhe	ValSerAsp	GluTyrPhePr	o SerGluPr
661		AATCAAATTA TTAGTTTAAT				

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	F16 7	CONTINUE	DII.			
+2	-	IleHisTyrAsı		ProGlnPhe	ThrGluAlaVa	l SerProSe
721		ATCCACTACA A				
+2	rValLeuPro	ProSerAlaLe	u ProLeuAsp	LeuLeuAsn	AsnAlaIleTh	r AlaPheSe
781		CCTTCAGCTT GGAAGTCGAA				
+2	rThrLeuGlu	AspLeulleAr	g TyrLeuGlu	ProGluArg	TrpGlnLeuAs	p LeuGluAs
841		GACCTTATTC ( CTGGAATAAG (				
. +2	pLeuTyrArg	ProThrTrpGl	n LeuLeuGly	LysAlaPhe	ValPheGlyAr	g LysSerAr
901		CCAACTTGGC A				
+2	gValValAsp	LeuAsnLeuLe	u ThrGluGlu	ValArgLeu	TyrSerCysTh	nr ProArgAs
961		CTGAACCTTC GACTTGGAAG				
+2	nPheSerVal	SerIleArgGl	u GluLeuLys	ArgThrAsp	ThrIlePheTi	p ProGlyCy
1021		TCCATAAGGG AGGTATTCCC			· ·	
+2	sLeuLeuVal	LysArgCysGl	y GlyAsnCys	AlaCysCys	LeuHisAsnCy	ys AsnGluCy
1081		AAACGCTGTG TTTGCGACAC				
+2	sGlnCysVal	ProSerLysVa	l ThrLysLys	TyrHisGlu	ValLeuGlnLe	eu ArgProLy
1141		CCAAGCAAAG GGTTCGTTTC				
+2	sThrGlyVal	ArgGlyLeuHi	s LysSerLe	ThrAspVal	AlaLeuGluH:	is HisGluGl
1201		AGGGGATTGC TCCCCTAACG				
+2		ValCysArgGl	-	_	•	
1261	GTGTGACTGT	GTGTGCAGAG CACACGTCTC	GGAGCACAGG	AGGATAGCCG		
1321		GTGCAGTGCA CACGTCACGT				
1381		GTTGTTTGCT				

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FIG. 7 (CONTINUED 2). 1441 AAGAGGAGAC ATCAAACAGA ATTAGGAGTT GTGCAACAGC TCTTTTGAGA GGAGGCCTAA TTCTCCTCTG TAGTTTGTCT TAATCCTCAA CACGTTGTCG AGAAAACTCT CCTCCGGATT 1501 AGGACAGGAG AAAAGGTCTT CAATCGTGGA AAGAAAATTA AATGTTGTAT TAAATAGATC TCCTGTCCTC TTTTCCAGAA GTTAGCACCT TTCTTTTAAT TTACAACATA ATTTATCTAG 1561 ACCAGCTAGT TTCAGAGTTA CCATGTACGT ATTCCACTAG CTGGGTTCTG TATTTCAGTT TGGTCGATCA AAGTCTCAAT GGTACATGCA TAAGGTGATC GACCCAAGAC ATAAAGTCAA 1621 CTTTCGATAC GGCTTAGGGT AATGTCAGTA CAGGAAAAAA ACTGTGCAAG TGAGCACCTG GAAAGCTATG CCGAATCCCA TTACAGTCAT GTCCTTTTTT TGACACGTTC ACTCGTGGAC 1681 ATTCCGTTGC CTTGGCTTAA CTCTAAAGCT CCATGTCCTG GGCCTAAAAT CGTATAAAAT TAAGGCAACG GAACCGAATT GAGATTTCGA GGTACAGGAC CCGGATTTTA GCATATTTTA 1741 CTGGATTTTT TTTTTTTTT TTGCGCATAT TCACATATGT AAACCAGAAC ATTCTATGTA GACCTAAAAA AAAAAAAAA AACGCGTATA AGTGTATACA TTTGGTCTTG TAAGATACAT 1801 CTACAAACCT GGTTTTTAAA AAGGAACTAT GTTGCTATGA ATTAAACTTG TGTCATGCTG GATGTTTGGA CCAAAAATTT TTCCTTGATA CAACGATACT TAATTTGAAC ACAGTACGAC 1861 ATAGGACAGA CTGGATTTTT CATATTTCTT ATTAAAATTT CTGCCATTTA GAAGAAGAGA TATCCTGTCT GACCTAAAAA GTATAAAGAA TAATTTTAAA GACGGTAAAT CTTCTTCTCT 1921 ACTACATTCA TGGTTTGGAA GAGATAAACC TGAAAAGAAG AGTGGCCTTA TCTTCACTTT TGATGTAAGT ACCAAACCTT CTCTATTTGG ACTTTTCTTC TCACCGGAAT AGAAGTGAAA 1981 ATCGATAGT CAGTTTATTT GTTTCATTGT GTACATTTTT ATATTCTCCT TTTGACATTA TAGCTATTCA GTCAAATAAA CAAAGTAACA CATGTAAAAA TATAAGAGGA AAACTGTAAT 2041 TAACTGTTGG CTTTTCTAAT CTTGTTAAAT ATATCTATTT TTACCAAAGG TATTTAATAT ATTGACAACC GAAAAGATTA GAACAATTTA TATAGATAAA AATGGTTTCC ATAAATTATA 2101 TCTTTTTAT GACAACTTAG ATCAACTATT TTTAGCTTGG TAAATTTTTC TAAACACAAT AGAAAAAATA CTGTTGAATC TAGTTGATAA AAATCGAACC ATTTAAAAAG ATTTGTGTTA 2161 TGTTATAGCC AGAGGAACAA AGATGATATA AAATATTGTT GCTCTGACAA AAATACATGT ACAATATCGG TCTCCTTGTT TCTACTATAT TTTATAACAA CGAGACTGTT TTTATGTACA 2221 ATTTCATTCT CGTATGGTGC TAGAGTTAGA TTAATCTGCA TTTTAAAAAA CTGAATTGGA TAAAGTAAGA GCATACCACG ATCTCAATCT AATTAGACGT AAAATTTTTT GACTTAACCT 2281 ATAGAATTGG TAAGTTGCAA AGACTTTTTG AAAATAATTA AATTATCATA TCTTCCATTC TATCTTAACC ATTCAACGTT TCTGAAAAAC TTTTATTAAT TTAATAGTAT AGAAGGTAAG 2341 CTGTTATTGG AGATGAAAAT AAAAAGCAAC TTATGAAAGT AGACATTCAG ATCCAGCCAT GACAATAACC TCTACTTTA TTTTTCGTTG AATACTTTCA TCTGTAAGTC TAGGTCGGTA 2401 TACTAACCTA TTCCTTTTTT GGGGAAATCT GAGCCTAGCT CAGAAAAACA TAAAGCACCT ATGATTGGAT AAGGAAAAAA CCCCTTTAGA CTCGGATCGA GTCTTTTTGT ATTTCGTGGA 2461 TGAAAAAGAC TTGGCAGCTT CCTGATAAAG CGTGCTGTGC TGTGCAGTAG GAACACATCC ACTITITCTG AACCGTCGAA GGACTATTTC GCACGACACG ACACGTCATC CTTGTGTAGG 2521 TATTTATTGT GATGTTGTGG TTTTATTATC TTAAACTCTG TTCCATACAC TTGTATAAAT ATAAATAACA CTACAACACC AAAATAATAG AATTTGAGAC AAGGTATGTG AACATATTTA

	F16.7	CONTINUE	ED 3).			
2581	ACATGGATAT	TTTTATGTAC	AGAAGTATGT	CTCTTAACCA	GTTCACTTAT	TGTACTCTGG
	TGTACCTATA	AAAATACATG	TCTTCATACA	GAGAATTGGT	CAAGTGAATA	ACATGAGACC
2641	CAATTTAAAA	GAAAATCAGT	AAAATATTTT	GCTTGTAAAA	TGCTTAATAT	CGTGCCTAGG
	GTTAAATTTT	CTTTTAGTCA	TTTTATAAAA	CGAACATTTT	ACGAATTATA	GCACGGATCC
2701	TTATGTGGTG	ACTATTTGAA	TCAAAAATGT	ATTGAATCAT	CAAATAAAAG	AATGTGGCTA
			AGTTTTTACA			
2761	TTTTGGGGAG	AAAATT				
	AAAACCCCTC	TTTTAA				

FIG. 8. Additional oligonucleotides used for amplification of entire coding region

5'-1	TTTGTTTAAACCTTGGGAAACTGG
5'-2	GTCCAGGTTTTGCTTTGATCC

# FIG. 9. DNA Sequence Of Clones 4 & 7, Identical Clones Containing The Entire Open Reading Frame

1		CCTTGGGAAA				
	AAACAAATTT	GGAACCCTTT	GACCAAGTCC	AGGTCCAAAA	CGAAACTAGG	AAAAGTTTTT
61		AGAAGAGGC				
	GACCTCTGTG	TCTTCTCCCG	AGATCCTTTT	TCAAAACCTA	CCCTAATACA	CCTTTGATGG
121		CTGCTGCCAG				
	GACGCTAAGA	GACGACGGTC	TCGTCCGAGC	CGCGAAGGTG	GGGTCACGTC	GGAAGGGGAC
181		AAGAGACTCG				
	CGCCACCACT	TTCTCTGAGC	CCTCAGCGAC	GAAGGTTTCA	CGGGCGCAC	TCACTCGAGA
+2	·, ·		_		LeuThrSerA	la LeuAlaGl
241	CACCCCAGTC	AGCCAAATGA	GCCTCTTCGG		CTGACATCTG	CCCTGGCCGG
		TCGGTTTACT				
+2	yGlnArgGln	GlyThrGlnA	la GluSerAsı	n LeuSerSer	LysPheGlnPl	ne SerSerAs
301	CCAGAGACAG	GGGACTCAGG	CGGAATCCAA	CCTGAGTAGT	AAATTCCAGT	TTTCCAGCAA
		CCCTGAGTCC				
+2	nLysGluGln	AsnGlyValG	ln AspProGlr	n HisGluArg	IleIleThrVa	al SerThrAs
361		AACGGAGTAC				
	GTTCCTTGTC	TTGCCTCATG	TTCTAGGAGT	CGTACTCTCT	TAATAATGAC	ACAGATGATT
+2	nGlySerIle	HisSerProA	rg PheProHis	ThrTyrPro	ArgAsnThrV	al LeuValTr
121		CACAGCCCAA				
	ACCTTCATAA	GTGTCGGGTT	CCAAAGGAGT	ATGAATAGGT	TCTTTATGCC	AGAACCATAC
+2	pArgLeuVal	AlaValGluG	lu AsnValTr	lleGlnLeu	ThrPheAspG	lu ArgPheGl
181		GCAGTAGAGG				
	CTCTAATCAT	CGTCATCTCC	TTTTACATAC	CTATGTTGAA	TGCAAACTAC	TTTCTAAACC
+2	yLeuGluAsp	ProGluAspAs		TyrAspPhe		lu GluProSe
541		CCAGAAGATG				
	CGAACTTCTG	GGTCTTCTAC	TGTATACGTT	CATACTAAAA	CATCTTCAAC	TCCTTGGGTC
+2	rAspGlyThr	IleLeuGlyA	rg TrpCysGly	y SerGlyThr	ValProGlyLy	ys GlnIleSe
501		ATATTAGGGC				
	ACTACCTTGA	TATAATCCCG	CGACCACACC	AAGACCATGA	CATGGTCCTT	TTGTCTAAAG
+2	rLysGlyAsn	GlnIleArgI:	le ArgPheVal	l SerAspGlu	TyrPheProS	er GluProGl
661		CAAATTAGGA				
		GTTTAATCCT				

	F/G. &	ALCONTINUED).	
+2		HisTyrAsnIle ValMetPro GlnPheThr GluAlaValSer ProSerV	'a
721	GTTCTGCATC CAAGACGTAG	CACTACAACA TTGTCATGCC ACAATTCACA GAAGCTGTGA GTCCTTCAGGTGATGTTGT AACAGTACGG TGTTAAGTGT CTTCGACACT CAGGAAGTC	T A
+2	lLeuProPro	SerAlaLeuPro LeuAspLeu LeuAsnAsn AlaIleThrAla PheSerT	'h
781		TCAGCTTTGC CACTGGACCT GCTTAATAAT GCTATAACTG CCTTTAGTA AGTCGAAACG GTGACCTGGA CGAATTATTA CGATATTGAC GGAAATCAT	
+2	rLeuGluAsp	LeuIleArgTyr LeuGluPro GluArgTrp GlnLeuAspLeu GluAspL	·е
841		CTTATTCGAT ATCTTGAACC AGAGAGATGG CAGTTGGACT TAGAAGATC GAATAAGCTA TAGAACTTGG TCTCTCTACC GTCAACCTGA ATCTTCTAG	
+2	uTyrArgPro	ThrTrpGlnLeu LeuGlyLys AlaPheVal PheGlyArgLys SerArgV	'a
901		ACTTGGCAAC TTCTTGGCAA GGCTTTTGTT TTTGGAAGAA AATCCAGAG TGAACCGTTG AAGAACCGTT CCGAAAACAA AAACCTTCTT TTAGGTCTC	
+2	lValAspLeu	AsnLeuLeuThr GluGluVal ArgLeuTyr SerCysThrPro ArgAsnP	'n
961		AACCTTCTAA CAGAGGAGGT AAGATTATAC AGCTGCACAC CTCGTAACT TTGGAAGATT GTCTCCTCCA TTCTAATATG TCGACGTGTG GAGCATTGA	
+2	eSerValSer	IleArgGluGlu LeuLysArg ThrAspThr IlePheTrpPro GlyCysL	,e
1021		ATAAGGGAAG AACTAAAGAG AACCGATACC ATTTTCTGGC CAGGTTGTC TATTCCCTTC TTGATTTCTC TTGGCTATGG TAAAAGACCG GTCCAACAG	
+2	uLeuValLys	ArgCysGlyGly AsnCysAla CysCysLeu HisAsnCysAsn GluCysG	1
1081		CGCTGTGGTG GGAACTGTGC CTGTTGTCTC CACAATTGCA ATGAATGTC GCGACACCAC CCTTGACACG GACAACAGAG GTGTTAACGT TACTTACAG	
+2	nCysValPro	SerLysValThr LysLysTyr HisGluVal LeuGlnLeuArg ProLysT	'h
1141		AGCAAAGTTA CTAAAAAATA CCACGAGGTC CTTCAGTTGA GACCAAAGA TCGTTTCAAT GATTTTTTAT GGTGCTCCAG GAAGTCAACT CTGGTTTCT	
+2	rGlyValArg	GlyLeuHisLys SerLeuThr AspValAla LeuGluHisHis GluGluC	<b>`</b> Y
1201		GGATTGCACA AATCACTCAC CGACGTGGCC CTGGAGCACC ATGAGGAGT CCTAACGTGT TTAGTGAGTG GCTGCACCGG GACCTCGTGG TACTCCTCA	
+2	sAspCysVal	CysArgGlySer ThrGlyGly	
1261		TGCAGAGGGA GCACAGGAGG ATAGCCGCAT CACCACCAGC AGCTCTTGCAGAGACGACCTCCCT CGTGTCCTCC TATCGGCGTA GTGGTGGTCG TCGAGAACG	
1321		CAGTGCAGTG GCTGATTCTA TTAGAGAACG TATGCGTTAT CTCCATCCTGTCACGTCAC	
1201			
1901		GTTTGCTTCA AGGACCTTTC ATCTTCAGGA TTTACAGTGC ATTCTGAAA CAAACGAAGT TCCTGGAAAG TAGAAGTCCT AAATGTCACG TAAGACTTT	
1441		AAACAGAATT AGGAGTTGTG CAA TTTGTCTTAA TCCTCAACAC GTT	

# FIG. 10. Predicted Full-length Polypeptide Sequence

1	MSLFGLLLLT	SALAGQRQGT	QAESNLSSKF	QF SSNKEQYG	VQDPQHERI
51	TVSTNGSIHS	PRFPHTYPRN	TVLVWRLVAV	EENVWIQLTF	DERFGLEDP
101	DDICKYDFVE	VEEPSDGTIL	GRWCGSGTVP	GKQISKGNQI	RIRFVSDEY
151	PSEPGFCIHY	NIVMPQFTEA	VSPSVLPPSA	LPLDLLNNAI	TAFSTLEDL
201	RYLEPERWQL	DLEDLYRPTW	QLLGKAFVFG	RKSRVVDLNL	LTEEVRLYS
251	TPRNFSVSIR	EELKRTDTIF	WPGCLLVKRC	GGNCACCLHN	CNECQCVPSI
301	VTKKYHEVLO	LRPKTGVRGL	HKSLTDVALE	HHEECDCVCR	GSTGG

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# FIG. 11. Alignment of VEGF-X with Other VEGFs

PLGF HUMAN :						: - : - : -
PLGF HUMAN :						: - : - : -
PLGF HUMAN :	DDICKYDFVEV		LGFFSVACSLL	AAALLPGPR	EAPAAAA EWVVVNV	: - : - : 30 : 10
PLGF_HUMAN : VEGB_HUMAN :	AFESGLDLSDA FMMLYVQLVQG		KDLEEQLRSVS	SVDELMTVL RAASSLEEL	MP  YPEYWKM LRITHSE	: 2 : -
VEGF_HUMAN : PLGF_HUMAN : VEGB_HUMAN : VEGC_HUMAN : VEGD_HUMAN : 990126vegx :	<ul><li>VMRLFPCFLQI</li><li>MSPLLRRI</li><li>YKCQLRKGGWQ</li></ul>	220 LLLYLHHAKWS( LAGLALPAVPPO LLAALLQLAPAO HNREQANLNSRI KSFTSMDSRSAS	AAPMAEGGGQN QWALSAGNGSS APVSQPDAPGH EETIKFAAAHY GHRSTRFAATFY	HHEVVKFMD EVEVVPFQE QRKVVSWID NTEILKSID DIETLKVID	-VWGRSY -VYTRAT NEWRKTQ EEWQRTQ	: 51 : 46 : 130 : 110
VEGF_HUMAN : PLGF_HUMAN : VEGB_HUMAN : VEGC_HUMAN : VEGD_HUMAN : 990126vegx :	CRALERLYDVV CQPREVVVPLT CMPREVCIDVC CSPRETCVEV	* 'QEYPDEIEYIFK 'SEYPSEVEHMFS 'VELMGTVAKQL' 'KEFGVATNTFFK 'SELGKSTNTFFK	PSCVSLLRCTG PSCVTVQRCGG PPCVSVYRCGG PPCVNVFRCGG	CCGD CCPD CCNS CCNE	300 EGLECVP ENLHCVP DGLECVP EGLOCMN ESLICMN NECOCVP	: 96 : 96 : 91 : 175 : 155 : 298

# FIG. 11 (CONTINUED).

		<u>*</u> 320 * 340 *		
VEGF_HUMAN	:	TEESNITMO MRIKPHOGOHIGEMSFLOWNKEERPKKDRAROEK	:	141
PLGF_HUMAN	:	VETANVTMOLLKURSGDRPSYVEUTFSOUVRCECRPLREKMKPER		141
VEGB_HUMAN	:	TEESNITMOTMRIKPHQGQHIGEMSFLQHNKCECRPKKDRARQEK VETANVTMQLLKIRSGDRPSYVELTFSQHVRCECRPLREKMKPER TGQHQVRMQILMIRYPSSQLGEMSLEEHSQCECRPKKKDSAVKP		135
VEGC_HUMAN		TSTSYLSKT FEITVPLSQGPKPVTISFANHTSCRCMSKLDVYRQVH	:	222
VEGD_HUMAN	:		•	
990126vegx	٠	TSTSYISKQIFEISVPLTSVPELVPVKVANHTGCKCLPTAPRHPYSI	:	202
990126Vegx	:	SKVTKKYHEVLQLRPKTGVRGLHKSLTDVALEHHEECDCVCRGSTGG	:	345
				•
		360 * 380 * 400	•	
THE THE THE SAN				
VEGF_HUMAN	:	KSVRGKGKGQKRKRKKSRYKSWSVP	:	166
PLGF_HUMAN	:		:	-
VEGB_HUMAN	:	DSPR	:	139
VEGC_HUMAN	:	SIIRRSLPATLPQCQAANKTCPTNYMWNNHICRCLAQEDFMFSSDAGDDS	:	272
VEGD_HUMAN	:	IRRSIQIPEEDRCSHSKKLCPIDMLWDSNKCKCVLQEENPLAGT	:	246
990126vegx	:		:	_
_				
		* 420 * 440 *		
VEGF_HUMAN	:		:	_
PLGF_HUMAN	:		:	_
VEGB_HUMAN	:			_
VEGC_HUMAN		TDGFHDICGPNKELDEETCQCVCRAGLRPASCGPHKELDRNSCQCVCKNK	:	322
VEGD_HUMAN	:	EDHSHLQEPALCGP	•	260
990126vegx	•	EDRSHUGEPALCGP	:	260
990126Vegx	•	***************************************	:	_
		460 * 480 * 500		
VEGF_HUMAN				206
PLGF_HUMAN		CCD TIDES	:	
	•		:	149
VEGB_HUMAN	:	PLCPRCTQHHQRPDPRTCRCRCRRRSFLRCQGRGLELNPD	:	179
VEGC_HUMAN	:	LFPSQCGANREFDENTCQCVCKRTCPRNQPLNPGKCACECTESPQKCLLK	:	372
VEGD_HUMAN	:	HMMFDEDRCECVCKTPCPKDLIQHPKNCSCFECKESLETCCQKHKLFHPD	:	310
990126vegx	:		:	-
		* 520 * 540 * TCRCDKPRR		
VEGF_HUMAN	:	TCRCDKPRR	:	215
PLGF_HUMAN	:		:	_
VEGB_HUMAN	:	TCRCRKLRR	•	188
VEGC_HUMAN	:	GKKFHHQTCSCYRRPCTNRQKACEPGFSYSEEVCRCVPSYWKRPQMS		419
VEGD HUMAN	:	TCSCEDRCPFHTRPCASGKTACAKHCRFPKEKRAAOGPHSRKNP	:	354
990126vegx	•	TOOCHDICE I HIN CABONIACAMICAL FARAMAQGEMSANIF	•	224
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	•	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	•	-

# FIG. 12. Variant Polypeptide Sequences

		* 20 * 40 *		
FL_seq	:	MSLFGLLLLTSALAGQRQGTQAESNLSSKFQFSSNKEQNGVQDPQHERII	:	50
clone41	:	MSLFGLLLLTSALAGQRQGTQAESNLSSKFQFSSNKEQNGVQDPQHERII	:	50
clone20	:	MSLFGLLLLTSALAGOROGTOAESNLSSKFOFSSNKEONGVODPOHERII	:	50
		60  * 80  * 100		
FL_seq	:	TVSTNGSIHSPRFPHTYPRNTVLVWRLVAVEENVWIQLTFDERFGLEDPE		100
clone41	:	TVSTNGSIHSPRFPHTYPRNTVLVWRLVAVEENVWIQLTFDERFGLEDPE	:	100
clone20	:	TVSTNGSIHSPRFPHTYPRNTVLVWRLVAVEENVWIQLTFDERFGLEDPE	:	100
EI com		* 120 * 140 *		
FL_seq clone41	•	DDICKYDFVEVEEPSDGTILGRWCGSGTVPGKQISKGNQIRIRFVSDEYF DDICKYDFVEVEEPSDGTILGRWCGSGTVPGKQISKGNQIRIRFVSDEYF	:	150
clone20	:	DDICKYDFVEVEEPSDGTILGRWCGSGTVPGKQISKGNQIRIRFVSDEYF	:	150 150
		TIAGO TATALQMBACIQABIVI BCOSMACCISCIAL VALVELLA CONTRACTOR CONTRAC	•	130
		160		
FL_seq	:	PSEPGFCIHYNIVMPQFTEAVSPSVLPPSALPLDLLNNAITAFSTLEDLI	:	200
clone41	:	PSEPSNRGGKIIOLHTS	:	167
clone20	:	PSEPGFCIHYN LVMPQFTEAVSPSVLPPSALPLDLLNNAITAFSTLEDLI	:	200
		* 220 * 240 *		
FL_seq	:	RYLEPERWQLDLEDLYRPTWQLLGKAFVFGRKSRVVDLNLLTEEVRLYSC		250
clone41	:	*	:	
clone20	:	RYLEPERWQLDLEDLYRPTWQLLGKAFVFGRKSRVVDLNLLTE	:	243
		260 * 280 * 300		
FL_seq		260 * 280 * 300 TPRNFSVSIREELKRTDTIFWPGCLLVKRCGGNCACCLHNCNECQCVPSK		2.00
clone41	:		:	3.00
clone20	:		•	_
			•	
TT		* 320 * 340		
FL_seq clone41	:	VTKKYHEVLQLRPKTGVRGLHKSLTDVALEHHEECDCVCRGSTGG : 345	,	
clone20	•	EVLQLRPKTGVRGLHKSLTDVALEHHEECDCVCRGSTGG : 282		
	•	- AAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA		

# F/G. 13. Primers for Expression of VEGF-X

#### E.coli expression of domain-

 vegx-6
 AATTGGATCCGAGAGTGGTGGATCTGAACC

 vegx-7
 AATTGGATCCGGGAAGAAAATCCAGAGTGG

 vegx-8
 GGTTGAATTCATTATTTTTTAGTAACTTTGCTTGGGACAC

 vegX-9
 AATTGAATTCATTATCCTCCTGTGCTCCCTC

Baculovirus/insect cell expression of full-length protein-

vegbac1

AATTGGATCCGGAGTCTCACCATCACCACCATCATGAATCCAACCTGAGTAGAATTCC

vegbac2 AATTGAATTCGCTATCCTCCTGTGCTCCCTCTGC

F16.14.

>3993180H1

LUNGNON03

INCYTE

CACAAATCACTCACCGACGTGGCCCTGGAGCACCATGAGGNGTGTGACTGTGTGTGCAGAGGGGGGAGCACAGGAGGATAGCC
GCATCACCACCAGCAGCTCTTGCCCAGAGCTGTGCAGTGCAGTGGCTGATTCTATTAGAGAACGTATGCGTTATCTCCAT
CCTTAATCTCAGTTGTTTGCTTCAAGGACCTTTCATCTTCAGGATTTACAGTGCATTCTGAAAGAGGAGACATCAAACAG
AATTAGGAGTTGTGCAACAGCTCTTTTGAGAGGAGGAGACATAAAGGACAGGAGAANAAGGTCTT

>3510192H1

CONCNOT01

INCYTE

>2559870H1

ADRETUT01

INCYTE

>3979767H1

LUNGTUT08

INCYTE

GGAGGATAGCCGCATCACCACCAGCAGCTCTTGCCCAGAGCTGTGCAGTGCAGTGGCTGATTCTATTAGAGAACGTATGC
GTTATCTCCATCCTTAATCTCAGTTGTTTGCTTCAAGGACCTTTCATCTTCAGGATTTACAGTGCATTCTGAAAGAAGAGAGACAGAACTAGCAACAGAACTAGCAACAGAACTAGAACAGAATTAGAGAGCAACAGCTCTTTTGAGAGAGGAGGCCTAAAGGACAGGAGAAAAAGGTCTTCAATCGTG
GAAAGAANATTAAATGTTGTATTAAATAGACACCAGCT

>3980011H1

LUNGTUT08

INCYTE

>4825396H1

BLADDIT01

INCYTE

GAGAACCGATACCATTTTCTGGCCAGGTTGTCTCCTGGTTAAACGCTGTGGTGGGAACTGTGCCTGTTGTCTCCACAATT GCAATGAATGTCAATGTGTCCCAAGCAAAGTTACTAAAAAATACCACGAGGTCCTTCAGTTGAGACCAAAGACCGGTGTC AGGGGATTGCACAAATCACTCACCGACGTGGCCCTGGAGCACCATGAGGAGTGTGACTGTGTGCAGAGGGAGCACAGG AGGATAGCCGCATCACCACCA

>3073703H1

BONEUNT01

INCYTE

AGAAAATCCAGAGTGGTGGATCTGAACCTTCTAACAGAGGAGGTAAGATTATACAGCTGCACACCTCGTAACTTCTCAGT GTCCATAAGGGAAGAACTAAAGAGAACCGATACCATTTTCTGGCCAGGTTGTCTCCTGGTTAAACGCTGTGGTGGGAACT GTGCCTGTTGTCTCCACAATTGCAATGAATGTCAATGTGTCCCAAGCAAAGTTACTAAAAAATACCACGAGGTCCTTCAG TTGAGACCAAAGACCGGTGTCAGGGGATTGCACAAATCA

>1302516H1

PLACNOT02

INCYTE

AGGAAATCAAATTAGGATAAGATTTGTATCTGATGAATATTTTCCTTCTGAACCTTCTAACAGAGGAGGTAAGATTATAC
AGCTGCACACCTCGTAACTTCTCAGTGTCCATAAGGGAAGAACTAAAGAGAACCGATACCATTTTCTGGCCAGGTTGTCT
CCTGGTTAAACGCTGTGGTGGGGAACTGTGCCTGTTGTCTCCCACAATTGCAATGTAATGTCAATGTCCCAAGCAAAGTT
ACTAAAAAAATACCACGAGGTCC

>3684109H1

HEAANOTO1 INCYTE

ATTTCATCTTCAGGATTTACAGTGCATTCTGAAANAGGAGAAATCAAACANAATTAGGAGTTGTGCAACAGCTCTTTTGA GAGGAGGCCTAAAGGACAGGAGAAAAGGTCTTCAATCGTGGAAANAAAATTAAATGTTGTATTAAATAGATCACCAGCTA GTTTCAGAGTTACCATGTACGTATTCCACTAGCTGGGTTCTGTATTTCAGTTCTTTCGATACGGCTTAGGGTAATGTCAG TACAGGAAAAAAACTGTGCAAGTGAGCACCTGATTCCGTTGCCTTGCTT

>4713188H1

BRAIHCT01

INCYTE

>458823H1

KERANOT01

INCYTE

ANGAGTTGCCCAGAGCTGTGCAGTGCAGTGGCTGATTCTATTAGAGAACGTATGCGTTATCTCCATCCTTAATCTCAGTT GTTTGNTTCAAGGACCTTTCATCTTCAGGATTTACAGTGCATTCTGAAAGAGGAGACATCAAACAGAATTAGGAGTTGTG CAACAGCTCTTTTGAGAGGAGGCCTAAAGGNCAGGAGAAAAGGTCTTCAATCGTGGAAAGAAAATTAAATGTTGTATTAA ATAGATC

>1303909H1

PLACNOT02

INCYTE

FIG. 14 (CONTINUED).

>2739211H1

OVARNOT09

INCYTE

GTGCATTCTGAAAGAGGAGACATCAAACAGAATTAGGAGTTGTGCAACAGCTCTTTTGAGAGGAGGGCCCTAAAGGACAGGA GAAAAGGTCTTCAATCGTGGAAAGAAAATTAAATGTTGTATTAAATAGATCACCAGCTAGTTTCAGAGTTACCATGTACG TATTCCACTAGCTGGGTTCTGTATTTCAGTTCTTTCGATACGGCTAAGGGTAATGTCAGTACAGGAAAAAAACTGTGCAA GTGAGCACCTGAT

>3325591H1

PTHYNOT03

INCYTE

>3733565H1

SMCCNOS01

INCYTE

CCTTAATCTCAGTTGTTTGCTTCAAGGACCTTTCATCTTCAGGATTTACAGTGCATTCTGNAAGANGAGACATCAAACAG
AATTAGGNGTTGTGCAAAAGCTCTTTTGAGAGGAGGGCCTAAAGGACAGGAGAAAAGGTCTNCAATCGTGGAAAGNAAATT
AAATGTTGTATNAAATNGATCACCAGCTAGTTTCAGAGTTACCATGTACGTATTCCACTAGCTGGGNCNGTATTCAGTCT
TTCGGAACGGCTTAGGGTAATGTCAGTACAGGANAAAAACTGTGCAGTGAG

>3554223H1

SYNONOT01

INCYTE

>4507477H1

OVARTDT01

INCYTE

 $\label{thm:control} \textbf{GGCTAGTTTCAGAGTTACCATTACGTATTCCACTAGCTGGGTTCTGTATTTCAGTTCTTTCGATACGGCTTAGGGTAAT\\ \textbf{GTCAGTACAGGAAAAAACTGTGCAAGTGAGCACCTGATTCCGTTGCCTTGACTCTAAAGCTCCATGTCCTGGGCC\\ \textbf{TAAAATCGTATAAAATCTGGA}$ 

>4163378H1

BRSTNOT32

INCYTE

AATAGATCACCAGCTAGTTTCAGAGTTACCATGTACGTATTCCACTAGCTGGGNTCTGTATTTCAGTTCCTTTCGATACG GCTTAGGGTAATGTCAGTACAGGAAAAAAGCTGTGCAAGTGAGCACCTGATTCCGTTGCCTTAACTCTAAAGCTCC ATGTCCTGGGCCTAAAATCGTATA F16.15.

>2054675H1

BEPINOT01

INCYTE

AAAGGAACTATGTTGCTATGAATTAAACTTGTGTCGTGCTGATAGGACAGACTGGATTTTTCATATTTCTTATTAAAATT
TCTGCCATTTAGAAGAAGAAGAACTACATTCATGGTTTGGAAGAGAGATAAACCTGAAAAGAAGAGGCCCTTATCTTCACTT
TATCGATAAGTCAGTTTATTTGTTTCATTGTGTACATTTTTATATTCTCCTTTTTGACATTATAACTGTTGGCTTTTCTAA
TCTTGTTAAATATCTATTTTTACCAAAGGTATTTAATATTCTTTTTTTA

>3993180H1

LUNGNON03

INCYTE

CACAAATCACTCACCGACGTGGCCCTGGAGCACCATGAGGNGTGTGACTGTGTGTGCAGAGGGGGGCACAGGAGGATAGCC GCATCACCACCAGCAGCTCTTGCCCAGAGCTGTGCAGTGCAGTGGCTGATTCTATTAGAGAACGTATGCGTTATCTCCAT CCTTAATCTCAGTTGTTTGCTTCAAGGACCTTTCATCTTCAGGATTTACAGTGCATTCTGAAAGAGGAGACATCAAACAG AATTAGGAGTTGTGCAACAGCTCTTTTGAGAGGAGGCTAAAGGACAGGAGAANAGGTCTT

>3510192H1

CONCNOT01

INCYTE

>4164633H1

BRSTNOT32

INCYTE

CTTGTTAAATATATCTATTTTTACCAAAGGTATTTAATATTCTTTANTTATGACAACTTAGATCAACTATTTTTAGCTTG GTAAATTTTTCTAAACACAATTGTTATAGCCAGAGGAACAAAGATGATATAAAAATATTGTTGCTCTGACAAAAATACATG TATTTCATTCTCGTATGGTGCTAGAGTTAGATTAATCTGCATTTTAAAAAAACTGAATTGGAATAGAATTGGTAAGTTGCA AAGACTTTTTGANAATAATTAAATTATCATATCTTCCATTCCTGTTATTGGGGGAGAAAAT

>2559870H1

ADRETUT01

INCYTE

>3817470H1

BONSTUT01

INCYTE

TTAAAAAGGAACTATGTTGCTATGAATTAAACTTGTGTCATGCTGATAGGACAGACTGGATTTTTCATATTTCTTATTAA
AATTTCTGCCATTTAGAAGAAGAAGAACTACATTCATGGTTTGGAAGAGAGATAAACCTGAAAAGAAGAGGCCTTATCTTC
ACTTTATCGATAAGTCAGTTTATTTGTTTCATTGTGTACATTTTTATATTCTCCTTTTGACATTATAACTGTTGGCTTTC
TAATCTGTTAAATATCTATTTTTACCAAAGGTATTTAATATTCTTT

>3979767H1

LUNGTUT08

INCYTE

GGAGGATAGCCGCATCACCAGCAGCTCTTGCCCAGAGCTGTGCAGTGCAGTGGCTGATTCTATTAGAGAACGTATGCGTTATCTCCATCCTTAATCTCAGTTGTTTGCTTCAAGGACCTTTCATCTTCAGGATTTACAGTGCATTCTGAAAGAGGAGACATCAAACAGACTTTTTGAGAGGAGGAGAAAAGGACAGGAGAAAAAGGTCTTCAATCGTGGAAAGAANATTAAATGTTGTATTAAATAGACACCAGCT

>3980011H1

LUNGTUT08

INCYTE

GGAGGATAGCCGCATCACCAGCAGCTCTTGCCCAGAGCTGTGCAGTGCAGTGGCTGATTCTATTAGAGAACGTATGC GTTATCTCCATCCTTAATCTCAGCTTGTTTGCCTTCAAGGACCTTTCATCTTCAGGATTTACATGCATTCTGAAAGAĞGAGA CATCAAACAGAATTAGGAGTTGTGCAACAGCTCTTTTGAGAGGAGGAGGCCTAAAGGACAGGAGAAAAGGTCTTCAATCGTGG AAAGAAAATTAAATGTTGTATTAAATAGATCACCA

>4825396H1

BLADDIT01

INCYTE

GAGAACCGATACCATTTTCTGGCCAGGTTGTCTCCTGGTTAAACGCTGTGGGGAACTGTGCCTGTTGTCTCCACAATT GCAATGAATGTCAATGTGTCCCAAGCAAAGTTACTAAAAAATACCACGAGGTCCTTCAGTTGAGACCAAAGACCGGTGTC AGGGGATTGCACAAATCACTCACCGACGTGGCCCTGGAGCACCATGAGGAGTGTGACTGTGTGCAGAGGGAGCACAGG AGGATAGCCGCATCACCACCA

>3073703H1

BONEUNT01

INCYTE

AGAAAATCCAGAGTGGTGGATCTGAACCTTCTAACAGAGGAGGTAAGATTATACAGCTGCACACCTCGTAACTTCTCAGT GTCCATAAGGGAAGAACTAAAGAGAACCGATACCATTTTCTGGCCAGGTTGTCTCCTGGTTAAACGCTGTGGTGGGAACT GTGCCTGTTGTCTCCACAATTGCAATGAATGTCAATGTGTCCCAAGCAAAGTTACTAAAAAAATACCACGAGGTCCTTCAG TTGAGACCAAAGACCGGTGTCAGGGGATTGCACAAATCA

>862169H1

BRAITUT03

INCYTE

AGATGATATAAAATATTGTTGCTCTGACAAAAATACATGTATTTCATTCTCGTATGGTGCTAGAGTTAGATTAATCTGCA TTTTAAAAAACTGAATTGGAATAGAATTGGTAAGTTGCAAAGACTTTTTGAAAATAATTAAATTATCATATCTTCCATTC CTGTTATTGGAGATGAAAATAAAAAGCAACTTATGAAAGTAGACATTCAGATCCAGCCATTACTAACCTATTCCTTTTTT GGGGAAATCTGAGCCTAGC

>4201385H1

BRAITUT29

INCYTE

TTTTTAAAAAGGAACTATGTTGCTATGAATTAAACTTGTGTCGTGCTGATAGGACAGACTGGATTTTTCATATTTCTTAT
TAAAATTTCTGCCATTTAGAAGAAGAAGAACTACATTCATGGTTTGGAAGAGATAAACCTGAAAAGAAGAGTGGCCTATCT
TCACTTTATCGATAAGTCAGTTTATTTGTTTCATTGTGTACATTTTTATATTCTCCTTTGACATATAACTGTTGGCTTTT

FIG. 15 (CONTINUED 1).

CTAATCTGTTAAATATATCTATTTTTACCAAAGGTATTTAATAT

>1302516H1

PLACNOT02

INCYTE

>3684109H1

HEAANOT01

INCYTE

ATTTCATCTTCAGGATTTACAGTGCATTCTGAAANAGGAGAAATCAAACANAATTAGGAGTTGTGCAACAGCTCTTTTGA GAGGAGGCCTAAAGGACAGGAGAAAAGGTCTTCAATCGTGGAAANAAAATTAAATGTTGTATTAAATAGATCACCAGCTA GTTTCAGAGTTACCATGTACGTATTCCACTAGCTGGGTTCTGTATTTCAGTTCTTTCGATACGGCTTAGGGTAATGTCAG TACAGGAAAAAAACTGTGCAAGTGAGCACCTGATTCCGTTGCCTTGCTT

>2549720H1

LUNGTUT06

INCYTE

>877279H1

LUNGAST01

INCYTE

CTTTTTTATGACAACTTAGATCAACTATTTTTAGCTTGGTAAATTTTTCTAAACACAATTGTTATAGCCAGAGGAACAAA GATGATATAAAATATTGTTGCTCTGACAAAAATACATGTATTTCATTCTCGTATGGTGCTAGAGTTAGATTAAATCTTGCAT TTTAAAAAAACTGAATTGGAATAGAATTGGTAAGTTGCAAAGGCTTTTTGAAAATAATTAAATTATCATATCTTCCATTCC TGTTATTGGNGG

>4713188H1

BRAIHCT01

INCYTE

>2171082H1

ENDCNOT03

INCYTE

AGATAAACCTGAAAAGAGAGGGCCTTATCTTCACTTTATCGATAAGTCAGTTTATTTTTTCATTGTGTACATTTTTA TATTCTCCTTTTGACATTATAACTGTTGGCTTTTCTAATCTTGTTAAATATATCTATTTTTACCAAAGGTATTTAATATT CTTTTTTATGACAACTTAGATCAACTATTTTTAGCTTGGTAAATTTTTCTAAACACAATTGTTATAGCCAGAGGAACAAA GATGA

>875860H1

LUNGAST01

INCYTE

>706168H1

SYNORAT04

INCYTE

GCTCATATTCACATATGTAAACCAGAACATTCTATGTACTACAAACCTGGTTTTTAAAAAGGANCTATGTTGCTATGAAT
TAAACTTGTGTCGTGCTGATAGGACAGACTGGATTTTTCATATTTCTTATTAAAATTTCTGCCATTTAGAAGAAGAAGAAC
TACATTCATGGTTTGGAAGAGAGATAAACCTGAAAAGAAGAGTGGCCTTATCTTCANTTTATCGATAAGTCAGTTTATTTGT
TTCA

>458823H1

KERANOT01

INCYTE

ANGAGTTGCCCAGAGCTGTGCAGTGCAGTGGCTGATTCTATTAGAGAACGTATGCGTTATCTCCATCCTTAATCTCAGTTGTTTGNTTCAAGGACCTTTCATCTTCAGGATTTACAGTGCATTCTGAAAGAGGAGACATCAAACAGAATTAGGAGTTGTGCAACAGCTCTTTTGAGAGGAGGAGGCCTAAAGGNCAGGAGAAAAGGTCTTCAATCGTGGAAAGAAAATTAAATGTTGTATTAAATGAGATC

>538436H1

LNODNOT02

INCYTE

AAAGATGATATAAAATATTGTTGCTCTGACAAAAATACATGTATTTCATTCTCGTATGGTGCTAGAGTTAGATTAATCTG CATTTTAAAAAACTGAATTGGAATAGAATTGGTAAGTTGCAAAGACTTTTTGAAAATAATTAAATTATCATATCTTCCAT TCCTGTTATTGGAGATGAAAATAAAAAGCAACTTATGAAAGTAGACATTCAGATCCAGCCATTACTAACCTAT

>1303909H1

PLACNOT02

INCYTE

>2739211H1

OVARNOT09

INCYTE

GTGCATTCTGAAAGAGAGACATCAAACAGAATTAGGAGTTGTGCAACAGCTCTTTTGAGAGGAGGCCTAAAGGACAGGA GAAAAGGTCTTCAATCGTGGAAAGAAAATTAAATGTTGTATTAAATAGATCACCAGCTAGTTTCAGAGTTACCATGTACG TATTCCACTAGCTGGGTTCTGTATTTCAGTTCTTTCGATACGGCTTAGGGTAATGTCAGTACAGGAAAAAAACTGTGCAA GTGAGCACCTGAT FIG. 15(continued 2).

>2550343H1

LUNGTUTOS

TNCYTE

TGTACATTTTATATTCTCCTTTTGACATTATAACTGTTGGCTTTTCNAATCTTGTTAAATATCTATTTTTACCAAAG GTATTTAATATTCTTTTTTATGACAACTTAGATCAACTATTTTTTAGCTTGGTAAATTTTTCTAAACACAATTGTTATAGC CAGAGGAACAAAGATGATATAAAATATTGTTGCTCTGACAAAAATACATGTATTTCATTCTCGTATGGTGCTA

>5321148H1

FIBPFEN06

INCYTE

>879495H1

THYRNOT02

INCYTE

ATTTCATTCTCGTATGGTGCTAGAGTTAGATTAATCTGCATTTTAAAAAACTGAATTGGAATAGAATTGGTAAGTTGCAAAGGCACTTTTTAAAAAACTGAATTGGAAAATAATTAAATTATCATATCTTCCATTCCTGTTATTGGAGATGAAAATAAAAAGCAACTTATGAAAAGTAAGCACCTTTTTTTGGGGAAATCTGAGCCTAGCTCAGAAAAAACATAAAGCACCTTGAAAAAA

>3325591H1

PTHYNOT03

INCYTE

>543890H1

OVARNOT02

INCYTE

TTTCTAAACACAATTGTTATAGCCAGAGGAACAAAGATGATATAAAATATTGTTGCTCTGACAAAAATACATGTATTTCA
TTCTCGTATGGTGCTAGAGTTAGATTAATCTGCATTTTAAAAAAACTGAATTGGNATAGAATTGGTAAGTTGCAAAGNCTT
TTTGAAAATAATTAAATTATCATATCTTCCATTCCTGTTATTGGAGGATGGAAAATAAAAAGCAACTTATGGAAAGTAGG
ACATTCAGATC

>3733565H1

SMCCNOS01

INCYTE

CCTTAATCTCAGTTGTTTGCTTCAAGGACCTTTCATCTTCAGGATTTACAGTGCATTCTGNAAGANGAGACATCAAACAG
AATTAGGNGTTGTGCAAAAGCTCTTTTGAGAGGAGGCCTAAAGGACAGGAGAAAAGGTCTNCAATCGTGGAAAGNAAATT
AAATGTTGTATNAAATNGATCACCAGCTAGTTTCAGAGTTACCATGTACGTATTCCACTAGCTGGGNCNGTATTCAGTCT
TTCGGAACGGCTTAGGGTAATGTCAGTACAGGANAAAAACTGTGCAGTGAG

>4641939H1

PROSTMT03

INCYTE

GTACTACAAACCTGGTTTTTAAAAAGGAACTATGTTGCTATGAATTAAACTTGTGTCCATGCTGATAGGACAGACTGGAT TTTNCATATTTCTTATTAAAATTTCTGCCATTTAGAAGAAGAGAACTACATTCATGGTTTGGNAGAGATAAACCTGAAAA GAAGAGTGGCCTTATCTTCACTTTATCGATAAGTCAGTTTATTTGTTTCATGTGTACATTTTTATATTCTCCTTTGACAT ATAACGTGGCTTT

>2007780H1

TESTNOT03

INCYTE

TTATATTCTCCTTTTGACATTATAACTGTTGGCTTTTCTAATCTTGTTAAATATATCTATTTTTACCAAAGGTATTTAAT
ATTCTTTTTTATGACAACTTAGATCAACTATTTTTAGCTTGGTAAATTTTTTCTAAACACAATTGTTATAGCCAGAGGAAC
AAAGATGATATAAAATATTGTTGCTCTGANAAAAATACATGTAT

>3085331H1

HEAONOT03

INCYTE

GCTCATATTCACATATGTAAACCAGAACATTCTATGTACTACAAACCTGGTTTTTAAAAAAGGAACTATTTGCTATGAATT
AAACTTGTGTCGTGCTGATAGGACAGACTGGNTTTTTCATATTTCTTATTANAATTTCTGCCATTAGAAGAAGAAGAACTA
CATTCATGGTTTTGGAAGAGAGATAAACCTGAAAAGAAGAGTGGCCTATTTCACTTTATCGATAAGTCAGT

>3414043H1

PTHYNOT04

INCYTE

GCTCATATTCACATATGTAAACCAGAACATTCTATGTACTACAAACCTGGTTTTTAAAAAGGAACTATGTTGCTATGAAT TAAACTTGTGTCGTGCTGATAGGACAGACTGGATTTTTCATATTTCTTATTAAAATTTCTGCCATTTAGAAGAAGAGAAC TACATTCATGGTTTGGAAGAGATAAACCTGAAA

>3705963H1

PENCNOT07

INCYTE

>5137051H1

OVARDIT04

INCYTE

>3554223H1

SYNONOT01

INCYTE

ATTAAATAGATCACCAGCTAGTTTCAGAGTTACCATGTACGTATTCCACTAGCTGGGTTCTGTATTTCAGTTCTTTCGAT ACGGCTTAGGGTAATGTCAGTACAGGAAAAAAACTGTGCAAGTGAGCACCTGATTCCGTTGCCTTAGCTTAACTCTAAAG

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### FIG. 15 (CONTINUED 3).

>4507477H1

OVARTDT01

INCYTE

 $\label{thm:control} \textbf{GGCTAGTTTCAGAGTTACCATGTATTCCACTAGCTGGGTTCTGTATTCAGTTCTTTCGATACGGCTTAGGGTAAT\\ \textbf{GTCAGTACAGGAAAAAAACTGTGCAAGTGAGCACCTGATTCCGTTGCCTTAACTCTAAAGCTCCATGTCCTGGGCC\\ \textbf{TAAAATCGTATAAAAATCTGGA}$ 

>1955646H1

CONNNOT01

INCYTE

TGGTAAGTTGCAAAGACTTTTTGAAAATAATTAAATTATCATATCTTCCATTCCTGTTATTGGAGATGAAAATAAAAAGC
AACTTATGAAAGTAGACATTCAGATCCAGCCATTACTAACCTATTCCTTTTTTTGGGGAAATCTGAGCCTAGCTCAGAAAA
ACATAAAGCACCTTGAAAAAGACTTGGCAGCTTCCTGATAAAGCGTGCTGTGCAGTAGGGAACACATCCTATTTA
TTGTGATGTTGTGGTTTTATATCCTAAACC

>4163378H1

BRSTNOT32

INCYTE

AATAGATCACCAGCTAGTTTCAGAGTTACCATGTACGTATTCCACTAGCTGGGNTCTGTATTTCAGTTCCTTTCGATACG GCTTAGGGTAATGTCAGTACAGGAAAAAAGCTGTGCAAGTGAGCACCTGATTCCGTTGCCTTAACTCTAAAGCTCC ATGTCCTGGGCCTAAAATCGTATA

>5095141H1

EPIMNON05

INCYTE

AGATAAACCTGAAAAGAGAGGGCCTTATNTTCACTTTATCGATAAGTCAGNTTATTTGTTTCATTGTGTACATTTNNA TATTCTCCTTTTGACATTATAACTGNTGGCTTTTCTAANCNTGTTAAATATATCTATTTTTACCAAAGGTATTTAATATT CTTT

>943826H1

ADRENOT03

INCYTE

>3451273H1

UTRSNON03

INCYTE

TTTTTTTTTTTGCTCATATTCACATATGTAAACCNGAACATCTATGTACNACAAACCTGGTTTTTAAAAAGGAACTATG TTGCTATGAATTAAACTTGTGTCGTGCTGATAGGACAGACTGGATTTTCANATTTCTTANTAANNTTTCTGCCATTTAG AAGA

>1402278H1

LATRTUT02

INCYTE

>4361191H1

SKIRNOT01

INCYTE

>1307017H1

PLACNOT02

INCYTE

>5032225H1

HEARFET03

INCYTE

>3732621H1

SMCCNOS01

INCYTE

>3530274H1

BLADNOT09

INCYTE

>3530249H1

BLADNOT09

INCYTE

# F1G. 16.

VEGFE1	AAAATGTATGGATACAACTTAC	22
VEGFE2	GTTTGATGAAAGATTTGGGCTTG	23
VEGFE3	TTTCTAAAGGAAATCAAATTAG	22
VEGFE4	GATAAGATTTGTATCTGATG	20
VEGFE5	GATGTCTCCTCTTTCAG	17
VEGFE6	GCACAACTCCTAATTCTG	18
VEGFE7	AGCACCTGATTCCGTTGC	19
VEGFE8	TAGTACATAGAATGTTCTGG	20
VEGFE9	AAGAGACATACTTCTGTAC	19
VEGFE10	CCAGGTACAATAAGTGAACTG	21

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F16.17 MNIFLL NLLTEEVRLY ]-----1 AGGAAATCAA ATTAGGATAA GATTTGTATC TGATGAATAT TTTCCTTCTG AACCTTCTAA CAGAGGAGGT AAGATTATAC TCCTTTAGTT TAATCCTATT CTAAACATAG ACTACTTATA AAAGGAAGAC TTGGAAGATT GTCTCCTCCA TTCTAATATG +3 S C T P R N F S V S I R E E L K R TDTIFWPGCL 81 AGCTGCACAC CTCGTAACTT CTCAGTGTCC ATAAGGGAAG AACTAAAGAG AACCGATACC ATTTTCTGGC CAGGTTGTCT TCGACGTGTG GAGCATTGAA GAGTCACAGG TATTCCCTTC TTGATTTCTC TTGGCTATGG TAAAAGACCG GTCCAACAGA \_\_\_\_\_\_ +3 L V K R C G G N C A C C L H N C N ECQCVPSKV 161 CCTGGTTAAA CGCTGTGGTG GGAACTGTGC CTGTTGTCTC CACAATTGCA ATGAATGTCA ATGTGTCCCA AGCAAAGTTA GGACCAATTT GCGACACCAC CCTTGACACG GACAACAGAG GTGTTAACGT TACTTACAGT TACACAGGGT TCGTTTCAAT -----+3 T K K Y H E V L Q L R P K T G V R G L H K S L T D V A +1 V S G D C T N H S P T W P 241 CTAAAAAATA CCACGAGGTC CTTCAGTTGA GACCAAAGAC CGGTGTCAGG GGATTGCACA AATCACTCAC CGACGTGGCC GATTTTTAT GGTGCTCCAG GAAGTCAACT CTGGTTTCTG GCCACAGTCC CCTAACGTGT TTAGTGAGTG GCTGCACCGG -----+3 L E H H E E C D C V C R G S T G G

### FIG. 17 (CONTINUED).

V Q R E H R R I A A S P P A A L A ]-----+1 WST MRSV TVC AEG AQED S R I T T S S S C 321 CTGGAGCACC ATGAGGAGTG TGACTGTGTG TGCAGAGGGA GCACAGGAGG ATAGCCGCAT CACCACCAGC AGCTCTTGCC GACCTCGTGG TACTCCTCAC ACTGACACAC ACGTCTCCCT CGTGTCCTCC TATCGGCGTA GTGGTGGTCG TCGAGAACGG +2 Q S C A V Q W L I L L E N V C V I SIL NLSC LLQ +1 P E L C S A V A D S I R E R M R Y L H P 401 CAGAGCTGTG CAGTGCAGTG GCTGATTCTA TTAGAGAACG TATGCGTTAT CTCCATCCTT AATCTCAGTT GTTTGCTTCA GTCTCGACAC GTCACGTCAC CGACTAAGAT AATCTCTTGC ATACGCAATA GAGGTAGGAA TTAGAGTCAA CAAACGAAGT

## +2 G P F I F R I Y S A F

- 481 AGGACCTTTC ATCTTCAGGA TTTACAGTGC ATTCTGAAAG AGGAGACATC AAACAGAATT AGGAGTTGTG CAACAGCTCT
  TCCTGGAAAG TAGAAGTCCT AAATGTCACG TAAGACTTTC TCCTCTGTAG
  TTTGTCTTAA TCCTCAACAC GTTGTCGAGA
- 561 TTTGAGAGGA GGCCTAAAGG ACAGGAGAAA AGGTCTTCAA TCGTGGAAAG AAAATTAAAT GTTGTATTAA ATAGATCACC
  AAACTCTCCT CCGGATTTCC TGTCCTCTTT TCCAGAAGTT AGCACCTTTC
  TTTTAATTTA CAACATAATT TATCTAGTGG
- 641 AGCTAGTTTC AGAGTTACCA TGTACGTATT CCACTAGCTG GGTTCTGTAT TTCAGTTCTT TCGATACGGC TTAGGGTAAT

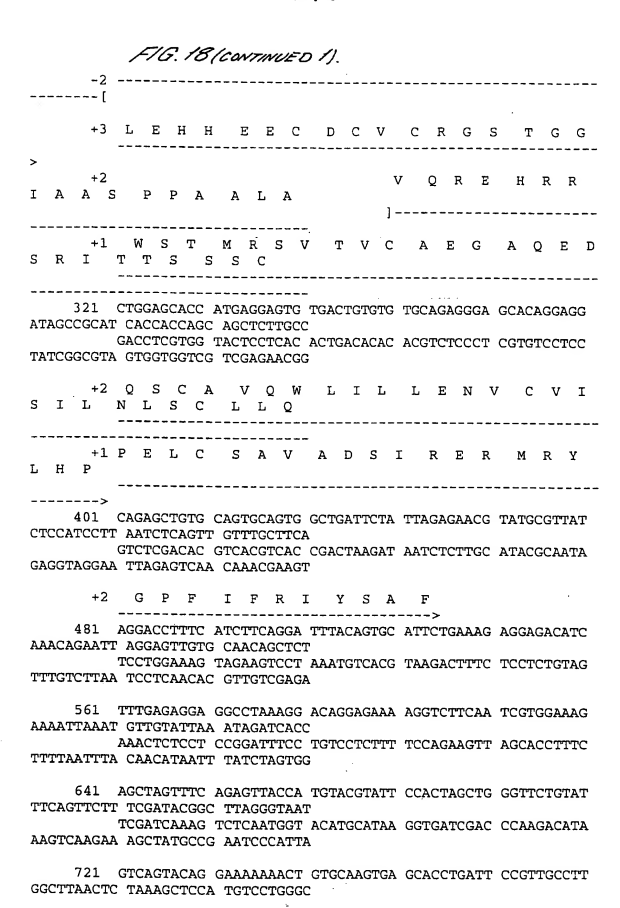
TCGATCAAAG TCTCAATGGT ACATGCATAA GGTGATCGAC CCAAGACATA AAGTCAAGAA AGCTATGCCG AATCCCATTA

- 721 GTCAGTACAG GAAAAAAACT GTGCAAGTGA GCACCTGATT CCGTTGCCTT
  GGCTTAACTC TAAAGCTCCA TGTCCTGGGC
- CAGTCATGTC CTTTTTTTGA CACGTTCACT CGTGGACTAA GGCAACGGAA CCGAATTGAG ATTTCGAGGT ACAGGACCCG
  - 801 CTAAAATCGT ATAAAATCTG GA GATTTTAGCA TATTTTAGAC CT

#### SUBSTITUTE SHEET (RULE 26)

FIG. 18.

NLLTEEV R-LY													
1 AGGAAATCAA ATTAGGATAA GATTTGTATC TGATGAATAT TTTCCAACCTTCTAA CAGAGGAGGT AAGATTATAC TCCTTTAGTT TAATCCTATT CTAAACATAG ACTACTTATA AAAGC													
TTGGAAGATT GTCTCCTCCA TTCTAATATG	SAAGAC												
+3 S C T P R N F S V S I R E E L T D T I F W P G C L													
81 AGCTGCACAC CTCGTAACTT CTCAGTGTCC ATAAGGGAAG AACTA AACCGATACC ATTTTCTGGC CAGGTTGTCT  TCGACGTGTG GAGCATTGAA GAGTCACAGG TATTCCCTTC TTGAT TTGGCTATGG TAAAAGACCG GTCCAACAGA -2	AAGAG												
+3 L V K R C G G N C A C C L H N E C Q C V P S K V													
161 CCTGGTTAAA CGCTGTGGTG GGAACTGTGC CTGTTGTCTC CACAATTGCA ATGAATGTCA ATGTGTCCCA AGCAAAGTTA GGACCAATTT GCGACACCAC CCTTGACACG GACAACAGAG GTGTTAACGT TACTTACAGT TACACAGGGT TCGTTTCAAT -2													
+3 T K K Y H E V L Q L R P K T G G L H K S L T D V A	V R												
	S G												
241 CTAAAAATA CCACGAGGTC CTTCAGTTGA GACCAAAGAC CGGTG GGATTGCACA AATCACTCAC CGACGTGGCC GATTTTTAT GGTGCTCCAG GAAGTCAACT CTGGTTTCTG GCCAC													



#### FIG. 18 (CONTINUED 2).

CAGTCATGTC CTTTTTTTGA CACGTTCACT CGTGGACTAA GGCAACGGAA CCGAATTGAG ATTTCGAGGT ACAGGACCCG

- 801 CTAAAATCGT ATAAAATCTG GATTTTTTTN TTTTTTTTTG CGCATATTCA CATATGTAAA CCAGAACATT CTATGTACTA
- GATTTTAGCA TATTTTAGAC CTAAAAAAAN AAAAAAAAAC GCGTATAAGT GTATACATTT GGTCTTGTAA GATACATGAT
- 881 CAAACCTGGT TTTTAAAAAG GAACTATGTT GCTATGAATT AAACTTGTGT CGTGCTGATA GGACAGACTG GATTTTTCAT
- GTTTGGACCA AAAATTTTTC CTTGATACAA CGATACTTAA TTTGAACACA GCACGACTAT CCTGTCTGAC CTAAAAAGTA
- -3 <-----
- 961 ATTTCTTATT AAAATTTCTG CCATTTAGAA GAAGAGAACT ACATTCATGG
  TTTGGAAGAG ATAAACCTGA AAAGAAGAGT

TAAAGAATAA TTTTAAAGAC GGTAAATCTT CTTCTCTGA TGTAAGTACC AAACCTTCTC TATTTGGACT TTTCTTCTCA

- -3 -----
- 1041 GGCCTTATCT TCACTTTATC GATAAGTCAG TTTATTTGTT TCATTGTGTA CATTTTATA TTCTCCTTTT GACATTATAA
- CCGGAATAGA AGTGAAATAG CTATTCAGTC AAATAAACAA AGTAACACAT GTAAAAATAT AAGAGGAAAA CTGTAATATT
  - -3 -----[
- 1121 CTGTTGGCTT TTCTAATCTT GTTAAATATA TCTATTTTTA CCAAAGGTAT TTAATATTCT TTTTTATGAC AACTTAGATC
- GACAACCGAA AAGATTAGAA CAATTTATAT AGATAAAAAT GGTTTCCATAAAATTATAAGA AAAAATACTG TTGAATCTAG
- 1201 AACTATTTT AGCTTGGTAA ATTTTTCTAA ACACAATTGT TATAGCCAGA GGAACAAAGA TGATATAAAA TATTGTTGCT
- TTGATAAAA TCGAACCATT TAAAAAGATT TGTGTTAACA ATATCGGTCT CCTTGTTTCT ACTATATTTT ATAACAACGA
- 1281 CTGACAAAA TACATGTATT TCATTCTCGT ATGGTGCTAG AGTTAGATTA ATCTGCATTT TAAAAAACTG AATTGGAATA
- GACTGTTTTT ATGTACATAA AGTAAGAGCA TACCACGATC TCAATCTAAT TAGACGTAAA ATTTTTTGAC TTAACCTTAT
- 1361 GAATTGGTAA GTTGCAAAGA CTTTTTGAAA ATAATTAAAT TATCATATCT TCCATTCCTG TTATTGGAGA TGAAAATAAA
- CTTAACCATT CAACGTTTCT GAAAAACTTT TATTAATTTA ATAGTATAGA AGGTAAGGAC AATAACCTCT ACTTTTATTT
- 1441 AAGCAACTTA TGAAAGTAGA CATTCAGATC CAGCCATTAC TAACCTATTC CTTTTTTGGG GAAATCTGAG CCTAGCTCAG
- TTCGTTGAAT ACTTTCATCT GTAAGTCTAG GTCGGTAATG ATTGGATAAG GAAAAAACCC CTTTAGACTC GGATCGAGTC

### FIG. 18 (CONTINUED 3).

- 1521 AAAAACATAA AGCACCTTGA AAAAGACTTG GCAGCTTCCT GATAAAGCGT GCTGTGCTGT GCAGTAGGAA CACATCCTAT
- 1601 TTATTGTGAT GTTGTGGTTT TATTATCTTA AACTCTGTTC CATACACTTG TATAAATACA TGGATATTTT TATGTACAGA
- AATAACACTA CAACACCAAA ATAATAGAAT TTGAGACAAG GTATGTGAAC ATATTTATGT ACCTATAAAA ATACATGTCT
  - AGTATGTCTC TTAACCAGTT CACTTATTGT ACCTGG TCATACAGAG AATTGGTCAA GTGAATAACA TGGACC

# FIG. 19. DNA and polypeptide sequence used for mammalian cell expression

- +1 m s l f g l l l l t s a l a g q r l GGATCCAAAA TGAGCCTCTT CGGGCTTCTC CTGCTGACAT CTGCCCTGGC CGGCCAGAGA
- +1 q g t q a E S N L S S K F Q F S S N K E 61 CAGGGGACTC AGGCGGAATC CAACCTGAGT AGTAAATTCC AGTTTTCCAG CAACAAGGAA
- +1 Q N G V Q D P Q H E R I I T V S T N G S 121 CAGAACGGAG TACAAGATCC TCAGCATGAG AGAATTATTA CTGTGTCTAC TAATGGAAGT
- +1 I H S P R F P H T Y P R N T V L V W R L
  181 ATTCACAGCC CAAGGTTTCC TCATACTTAT CCAAGAAATA CGGTCTTGGT ATGGAGATTA
- +1 V A V E E N V W I Q L T F D E R F G L E 241 GTAGCAGTAG AGGARAATGT ATGGATACAA CTTACGTTTG ATGALAGATT TGGGCTTGAA
- +1 D P E D D I C K Y D F V E V E E P S D G 301 GACCCAGAAG ATGACATATG CAAGTATGAT TTTGTAGAAG TTGAGGAACC CAGTGATGGA
- +1 T I L G R W C G S G T V P G K Q I S K G 361 ACTATATAG GGCGCTGGTG TGGTTCTGGT ACTGTACCAG GAAAACAGAT TTCTAAAGGA
- +1 N Q I R I R F V S D E Y F P S E P G F C
- +1 I H Y N I V M P Q F T E A V S P S V L P 481 ATCCACTACA ACATTGTCAT GCCACAATTC ACAGAAGCTG TGAGTCCTTC AGTGCTACCC
- +1 P S A L P L D L L N N A I T A F S T L E 541 CCTTCAGCTT TGCCACTGGA CCTGCTTAAT AATGCTATAA CTGCCTTTAG TACCTTGGAA
- +1 D L I R Y L E P E R W Q L D L E D L Y R 601 GACCTTATTC GATATCTTGA ACCAGAGAGA TGGCAGTTGG ACTTAGAAGA TCTATATAGG
- +1 P T W Q L L G K A F V F G R K S R V V D
  661 CCAACTTGGC AACTTCTTGG CAAGGCTTTT GTTTTTGGAA GAAAATCCAG AGTGGTGGAT
- +1 L N L L T E E V R L Y S C T F R N F S V
  721 CTGAACCTTC TAACAGAGGA GGTAAGATTA TACAGCTGCA CACCTCGTAA CTTCTCAGTG
- +1 S I R E E L K R T D T I F W P G C L L V 781 TCCATAAGGG AAGAACTAAA GAGAACCGAT ACCATTTTCT GGCCAGGTTG TCTCCTGGTT
- +1 K R C G G N C A C C L H N C N E C Q C V 841 AAACGCTGTG GTGGGAACTG TGCCTGTTGT CTCCACAATT GCAATGAATG TCAATGTGTC
- +1 P S K V T K K Y H E V L Q L R P K T G V 901 CCAAGCAAAG TTACTAAAAA ATACCACGAG GTCCTTCAGT TGAGACCAAA GACCGGTGTC
- +1 R G L H K S L T D V A L E H H E E C D C 961 AGGGGATTGC ACAAATCACT CACCGACGTG GCCCTGGAGC ACCATGAGGA GTGTGACTGT
- +1 V C R G S T G G <u>S R G P F E G K P I P N</u>
  1021 GTGTGCAGAG GGAGCACAGG AGGATCTAGA GGGCCCTTEG AAGGTAAGCC TATCCCTAAC

#### FIG. 20. DNA and polypeptide sequence used for baculovirus/insect cell expression

- 1 GAATTCAAAG GCCTGTATTT TACTGTTTTC GTAACAGTTT TGTAATAAAA AAACCTATAA
- +3 m k f l v n v a l v f m v v y i s y i 61 ATATGAAATT CTTAGTCAAC GTTGCCCTTG TTTTTATGGT CGTATACATT TCTTACATCT
- +3 Y a D P E S H H H H H E S N L S S K F
  121 ATGCGGATCC GGAGTCTCAC CATCACCACC ATCATGAATC CAACCTGAGT AGTAAATTCC
- +3 Q F S S N K E Q N G V Q D P Q H E R I I 181 AGTTTCCAG CAACAAGGAA CAGAACGGAG TACAAGATCC TCAGCATGAG AGAATTATTA
- +3 T V S T N G S I H S P R F P H T Y P R N
  241 CTGTGTCTAC TAATGGAAGT ATTCACAGCC CAAGGTTTCC TCATACTTAT CCAAGAAATA
- +3 T V L V W R L V A V E E N V W I Q L T F 301 CGGTCTTGGT ATCCAGATTA GTAGCAGTAG AGGAAAATGT ATGGATACAA CTTACGTTTG
- +3 D E R F G L E D P E D D I C K Y D F V E 361 ATGAAAGATT TGGGCTTGAA GACCCAGAAG ATGACATATG CAAGTATGAT TTTGTAGAAG
- +3 V E E P S D G T I L G R W C G S G T V P
  421 TTGAGGAACC CAGTGATGGA ACTATATTAG GGCGCTGGTG TGGTTCTGGT ACTGTACCAG
- +3 G K Q I S K G N Q I R I R F V S D E Y F 481 GAAAACAGAT TTCTAAAGGA AATCAAATTA GGATAAGATT TGTATCTGAT GAATATTTTC
- +3 P S E P G F C I H Y N I V M P Q F T E A 541 CTTCTGAACC AGGGTTCTGC ATCCACTACA ACATTGTCAT GCCACAATTC ACAGAAGCTG
- +3 V S P S V L P P S A L P L D L L N N A I 601 TGAGTCCTTC AGGGCTACCC CCTTCAGCTT TGCCACTGGA CCTGCTTAAT AATGCTATAA
- +3 T A F S T L E D L I R Y L E P E R W Q L 661 CTGCCTTTAG TACCTTGGAA GACCTTATC GATATCTTGA ACCAGAGAGA TGGCAGTTGG
- +3 D L E D L Y R P T W Q L L G K A F V F G
  721 ACTTAGAAGA TCTATATAGG CCAACTTGGC AACTTCTTGG CAAGGCTTTT GTTTTTGGAA
- +3 R K S R V V D L N L L T E E V R L Y S C 781 GAAAATCCAG AGTGGTGGAT CTGAACCTTC TAACAGAGGA GGTAAGATTA TACAGCTGCA
- +3 T P R N F S V S I R E E L K R T D T I F 841 CACCTCGTAA CTTCTCAGTG TCCATAAGGG AAGAACTAAA GAGAACCGAT ACCATTTTCT
- +3 W P G C L L V K R C G G N C A C C L H N
  901 GGCCAGGTTG TCTCCTGGTT AAACGCTGTG GTGGGAACTG TGCCTGTTGT CTCCACAATT
- +3 C N E C Q C V P S K V T K K Y H E V L Q
  961 GCAATGAATG TCAATGTGTC CCAAGCAAAG TTACTAAAAA ATACCACGAG GTCCTTCAGT
- +3 L R P K T G V R G L H K S L T D V A L E
  1021 TGAGACCAAA GACCGGTGTC AGGGGATTGC ACAAATCACT CACCGACGTG GCCCTGGAGC
- +3 H H E E S D C V C R G S T G G
  1081 ACCATGAGA GIGIGACTGT GTGTGCAGAG AGGACACAG AGGATAGCTC TAGA

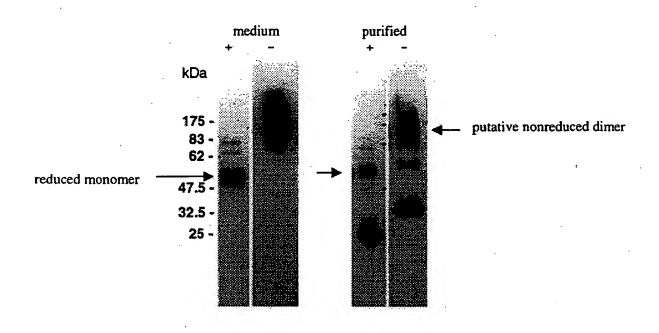
WO 00/37641 PCT/US99/30503

FIG. 21. DNA and polypeptide sequence used for E.coli expression

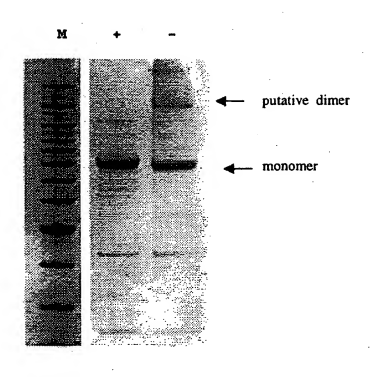
+3	OTN	S S S	NNNN	<u> </u>	N N N	L G I
1	CGCAGACTAA	TTCGAGCTCG	AACAACAACA	ACAATAACAA	TAACAACAAC	CTCGGGATCG
<b></b> 3	FGPT	9 6 6	E S N L		FOF	SSN
			GAATCCAACC			
		_	D P Q H			
121	AGGAACAGAA	CGGAGTACAA	GATCCTCAGC	ATGAGAGAAT	TATTACTGTG	TCTACTAATG
+3	G S I H	SPR	F P H T	YPR	N T V	L V W
			TTTCCTCATA			
_			·			
			N V W I			
	0					
			тску			
301	TTGAAGACCC	AGAAGATGAC	ATATGCAAGT	ATGATTTTGT	AGAAGTTGAG	GAACCCAGTG
+3	DGTI	ī. G. R	WCGS	G T V	рск	0 T S
			TGGTGTGGTT			
	_		R F V S AGATTTGTAT			
421	AAGGAAAICA	AAITAGGATA	AGATTTGTAT	CIGAIGAAIA	TTTTCCTTCT	GAACCAGGGT
+3	F C I H	YNI	V M P Q	FTE	A V S	P S V
481	TCTGCATCCA	CTACAACATT	GTCATGCCAC	AATTCACAGA	AGCTGTGAGT	CCTTCAGTGC
. 2	מ מ מ	3 T D	LDLL	N N A	T T A	E C T
			CTGGACCTGC			
			L E P E			
601	TGGAAGACCT	TATTCGATAT	CTTGAACCAG	AGAGATGGCA	GTTGGACTTA	GAAGATCTAT
+3	Y R P T	W Q L	LGKA	FVF	G R K	s R V .
661	ATAGGCCAAC	TTGGCAACTT	CTTGGCAAGG	CTTTTGTTTT	TGGAAGAAAA	TCCAGAGTGG
. 3	V D I M	7 7 m	E E V R	7 V C	C 17 D	D W E
	_		GAGGAGGTAA			
			LKRT			
781	CAGTGTCCAT	AAGGGAAGAA	CTAAAGAGAA	CCGATACCAT	TTTCTGGCCA	GGTTGTCTCC
+ 3	LVKR	CGG	N C A C	сгн	N C N	E C Q
			AACTGTGCCT			
			K K Y H			
301	J.G.CCCAAG	Crance INCI	- STATEMENT ACC	right i Weigh	·	UUUMUMUUUU
+3	G V R G	L H K	S L T D	V A L	Е Н Н	E E C
961	GTGTCAGGGG	ATTGCACAAA	TCACTCACCG	ACGTGGCCCT	GGAGCACCAT	GAGGAGTGTG
. 7		2 6 6	т		и и *	
			ACAGGAGGAC			TCTAGAGTCG
1081	ACCTGCAGGC	AAGCTT	•			•

# FIG. 22. Disulphide-linked dimerisation of VEGF-X

#### (A) Mammalian cell expression



#### (B) E.coli expression



# FIG. 23. Glycosylation of VEGF-X

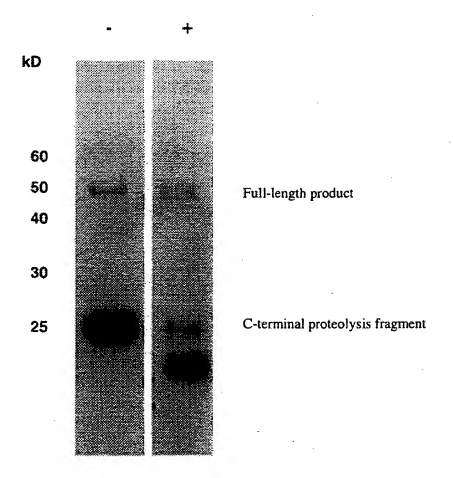


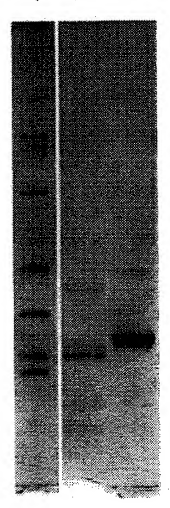
FIG. 24.

DNA and polypeptide sequence used for E.coli expression of the PDGF-like domain

+3						M	R	G	s	н	н	н	н	H	н	G	М	_ A	S	М	
1	AA	GGA	GAT	ΑT	ACAT	ATG	CGG	GGT	TCI	CAT	c .	ATCAT	CAT	CA	TCAT	GGT	ATG	GCT	AGC	ATGA	
+3	<u>T_</u>	G	G	0	0	M	G	R	D	L	Y	Q	_D	D	D	K	D	P	G	R	
61	1 CTGGTGGACA			GCAA	ATG	GGT	CGG	GAT	TCTGT		ACGAC	GAT	ĠΑ	CGAT	'AAG	GAT	CCGGGAAGA				
+3	K	s	R	٧	v	D	L	N	L	L	T	E	E	V	R	L	Y	s	С	T	
121	AA	TCC	AGA	ЗT	GGTG	GAT	CTG	AAC	CTI	CTA	A	CAGAG	GAG	GT	AAGA	ATT.	TAC	AGCTGCAC			
			. •																		
+3	P	R	N	F	S	V	S	I	R	E	E	L	K	R	T	D	T	I	F	W	
181	CT	CGT	AAC.	ΓT	CTCA	GTG	TCC	ATA	AGG	GAA	G.	AACTA	AAG	AG	AACCGATACC			ATTTTCTGG			
+3	₽	G	C	L	L	V	K	R	C	G	G	N	С	A	С	С	L	H	N	С	
241	CA	GGT"	rgr(	CT	CCTG	GTT	'AAA	CGC	TGI	GGT	G	GGAAC	TGT	'GC	CTGT	TGT	CTC	CAC	TA_5	TGC	
				_								K									
301	AT	GAA'	TGT(	CA	ATGI	'GTC	CCA	AGCAAAGTTA CTAAAAAATA						CCAC	GAG	GTC	CTTCAGTTG				
_	_	_																			
												S									
361	GA	CCA	AAG	AC	CGGI	GTC	AGG	GGATTGCACA AATCACTCAC						AC	CGAC	:GTG	GCC	CIG	GAG	CAC	
_		_	_	_	_	_		_	_	_	_	_	_	_							
				_	_	_		_		-	_	T	_	_							
421	AT	GAG	JAG'	ľĠ	TGAC	TGT	GTG	TGC	AGA	.GGG	A	GCACA	LGGA	.GG	ATAA	TGA	ATT	CGA	AGC	TTG	
481	TC	الحالات	CIG	_T	AACA	AAG	CCC														

F/G. 25. Expression of PDGF domain in E.coli

1 2 3



# FIG. 26.

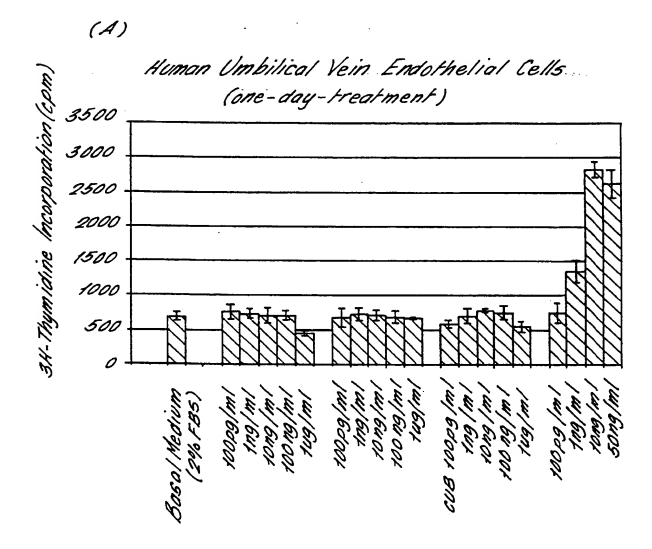
# DNA and polypeptide sequence used for E.coli expression of the CUB-like domain

+2		M	Α_	M	ם	I	G	I	_N	_s	D	P	E	S	н	н	н	H	н	н	
													TCTCACCATC ACCACCATCA								
			• 0																		
	Ε																				
61	TGAA	TCC	AAC	CTG	AGI	AGT	A 1	AATTO	CAG	TT	TTC	AGC	AAC	AAG	AAGGAACAGA ACGGAGTAG						
<b>+</b> 2	n	ъ	^	22	-	-	<b>T</b>	-	~	,,	_		.,	_	_	_		_	_		
	D AGAT																				
121	AGAT	-C-1	CAG	CAI	GAC	MOM	•	IIAII	ACI	GI	GICI	AUI	AAT	GGA	AGT	ATT	C A	CAG	CCA	LAG	
+2	F	P	н	Т	Y	P	R	N	т	v	L	. v	W	R	Ť.	v	Δ	v	F	F	
	GTTT																				
+2	N	v	W	I	Q	L	T	F	D	E	R	F	G	L	E	D	P	Ε	D	D	
241	AAAT	GTA'	TGG	ATA	CAA	CTT	١ (	GTTI	GAT	GA	AAGATTTGGG CTTGAAGACC CAGAA						GAT	'GA			
										•											
+2	I	С	K	Y	D	F	V	E	V	E	E	P	S	D	G	T	I	L	G	R	
301	CATA	TGC	AAG	TAT	GAT	TTT	; 7	ragaa	GTT	GA	GGAACCCAGT GATGGAACTA TATTAGGGC							CG			
	W																				
361	CTGG'	IGT(	GGT	TCT	GGT	ACTO	7	TACCA	GGA	AA	ACAG	ATT	TCT	AAA	GGA.	AAT	C A	AATI	'AGG	AT	
_	_	_		_	_	_		_													
	R																				
421	AAGA	TTT(	JTA	TCT	GAT	GAAT	: <i>F</i>	ATTTT	'CCT	TC	TGAA	CCA	.GGG	TTC	TGC	ATC	C A	CTAC	AAC	ΑT	
	v	M	ם	_		T	-	*	**												
				_						<b></b>	cmcc		mca		~~						
-201	TGTC	-10		بربي		r.CAC		MOCI	GIG	IA	GICE	MGC	100	GIC	GAC	AAG	C T	TGCG	GCC	.GC	
541	ACTC	GAG	CAC																		

# F/G. 27. Expression of the CUB domain in E.coli

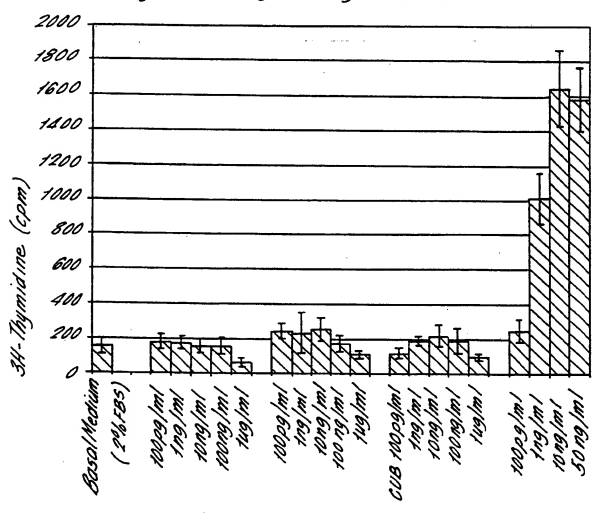


FIG. 28. The Effect of Truncated VEGF-X (CUB domain) on HUVEC Proliferation.



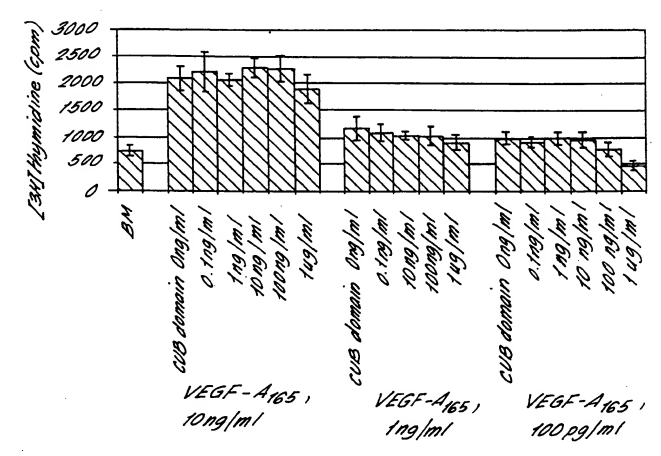
### F1G. 28(CONTINUED 1).

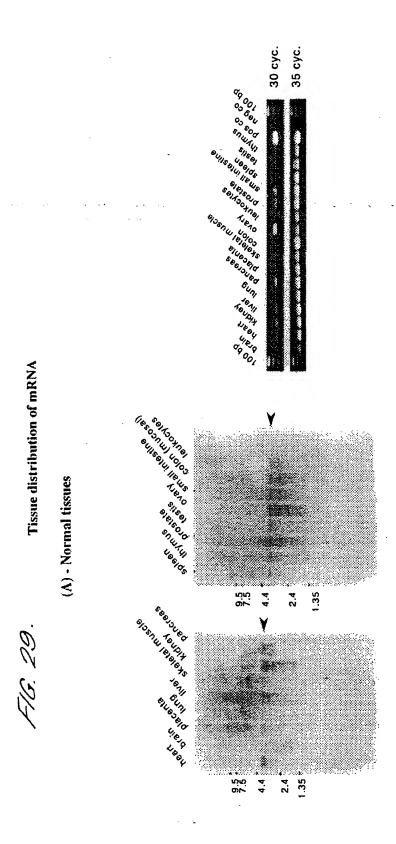
(8)
Human Umbilical Vein Endothelial Cells (24-hourstorving Followed by one-day-treatment)



### FIG. 28 (CONTINUED 2).

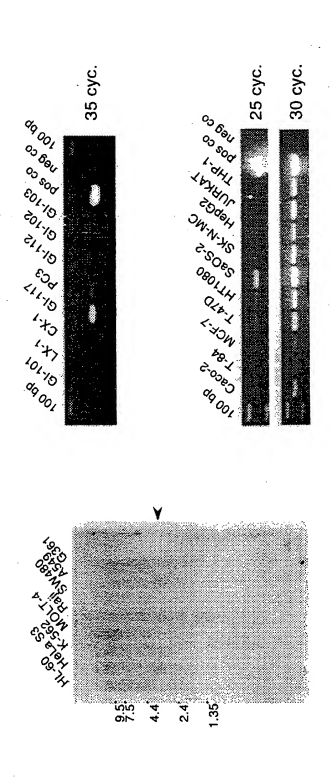
(C)
The effect of VEGF-A<sub>165</sub> and VEGF-X CUB domain on the proliferation of HUVEC (two-day-treatment).





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FIG. 29 (covernue D). (B)- Tumour tissue and cell lines



F1G.30

#### Partial intron/exon structure of the VEGF-X gene

#### (A) - Genomic DNA sequences of 2 exons determined by sequencing

aaagccagtcatagacattcgttgatttttaaaagtggcttactcttattccctttcagGTCCTTCAGTTGAGACCAAAGACCGGT GTCAGGGGATTGCACAAATCACTCACCGACGTGGCCCTGGAGCACCATGAGGAGTGTGACTGTGTGCAGAGGGAGCACAGGAGG ATAGCCGCATCACCACCAGCAGCTCTTGCCCAGAGCTGTGCAGTGCAGTGGCTGATTCTATTAGAGAACGTATGCGTTATCTCCAT AAATAGATCACCAGCTAGTTTCAGAGTTACCATGTACGTATTCCACTAGCTGGGTTCTGTATTTCAGTTCTTTCGATACGGCTTAG GGTAATGTCAGTACAGGAAAAAACTGTGCAAGTGAGCACCTGATTCCGTTGCCTTGGCTTAACTCTAAAGCTCCATGTCCTGGGC TGGTTTTTAAAAAGGAACTATGTTGCTATGAATTAAACTTGTGTCATGCTGATAGGACAGACTGGATTTTTCATATTTCTTATTAA AATTTCTGCCATTTAGAAGAAGAACTACATTCATGGTTTGGAAGAGATAAACCTGAAAAGAAGAGTGGCCTTATCTTCACTTTA TCGATAAGTCAGTTTATTTGTTTCATTGTGTACATTTTTATATTCTCCTTTTGACATTATAACTGTTGGCTTTTCTAATCTTGTTA AATATATCTATTTTTACCAAAGGTATTTAATATTCTTTTTTATGACAACTTAGATCAACTATTTTTAGCTTGGTAAATTTTTCTAA ACACAATTGTTATAGCCAGAGGAACAAAGATGATATAAAATATTGTTGCTCTGACAAAAATACATGTATTTCATTCTCGTATGGTG CTAGAGTTAGATTAATCTGCATTTTAAAAAACTGAATTGGAATAGAATTGGTAAGTTGCAAAGACTTTTTGAAAATAATTAAATTA TCATATCTTCCATTCCTGTTATTGGAGATGAAAATAAAAAGCAACTTATGAAAGTAGACATTCAGATCCAGCCATTACTAACCTAT TCCTTTTTTGGGGAAATCTGAGCCTAGCTCAGAAAAACATAAAGCACCTTGAAAAAGACTTGGCAGCTTCCTGATAAAGCGTGCTG TGCTGTGCAGTAGGAACACATCCTATTTATTGTGATGTTGTGGTTTTATTATCTTAAACTCTGTTCCATACACTTGTATAAATACA TGGATATTTTATGTACAGAAGTATGTCTCTTAACCAGTTCACTTATTGTACTCTGGCAATTTAAAAGAAAATCAGTAAAATATTT TGCTTGTAAAATGCTTAATATCGTGCCTAGGTTATGTGGTGACTATTTGAATCAAAAATGTATTGAATCATCAAATAAAAGAATGT GGCTATTTTGGGGAGAAAATTatgtgtgtgtgtgttcaagatttatttcttggactctgagaaaatgaaagataaa

#### FIG. 30 (CONTINUED 1).

#### (B) - Location of splice sites within the cDNA sequence

1	GAA:	TTC	GCCC	TTT	CTI	TAA	ACC	TTG	GGA	4 (	TGGT	TCAG	G	TCCA	GGT	TTT	GCT	TTG	ATCC
61	TTT	CA	AAAA	CTG	AGA	CAC	AGA	AGA	GGG	<b>=</b> 1	CTAG	GAAA	LA	AGTT	TTG	GAT	GGG.	ATT!	atgt
121	GGA	AC.	TACC	CTG	GAI	TCT	CTG	CTG	CCAC	3 A	AGCAG	GCTC	G	GCGC	TTC	CAC	CCC	AGTO	GCAG
181	CCT	rcc	CTG	GCGC	TGG	TGA	AAG	AGA	CTC	3 6	GAGT	CGC1	C	CTTC	CAA.	AGT	GCC	CGC	CGTG
+3									М	s	L	F	G	L	L	L	L	T	s
241		iag	CTCT	CACO	CCA	GTC													
	A 1		a	0	P	0	Ġ	T	0	Δ	E	s	N	T.	s	s	X	F	0
301																			
. 2	r (		e at	2	 E		AT	G	v	0	D.	D	<u> </u>	น	F	P	T	т	т
361																			
	17 (	. ,	T 37	_	c	_	ŧ	c	D	Ð	F	D	Ľ	T	v	Ð	D	N	$\boldsymbol{\tau}$
421																			
. 7	١,	. 1	, W	2	7	7.7	a	1/	F	F	N	v	w	7	0	r.	T	F	ת
481																			
+3	E F	۱ ۶	F G	L	E	D	₽	E	D	D	I	С	K	Y	D	F	V	E	V
541	AAA	JAT:	TTGG	GCTT	rgaa	GAC	CCA	GAA	GAT(	3 2	CATA	TGC	lA	GTAT	GAT	TTT	GTA	GAX(	STIG
+3	E I	E 1	P S	D	G	T	·I	L	G	R	W	С	G	s	G	T	v	Þ	G
601	AGG	AC	CCAG	TGAT	rgga	ACT	ATA	TTA	GGG	c	CTGG	TGTC	G	TTCT	GGT.	ACT	GTA	CCA	ggaa
											R								
661	AAC	AGA:	TTTC	TAA	<i><b>IGGA</b></i>	AAT	CAA	ATT	AGG	A 7	raaga	TTTC	T	ATCT	GAT	GAA	TAT	777	CCTT
		_ ,	- 1-	_	_			•	17	_	v	<b>W</b>	ъ	^	E	T	-	n.	11
721	CTG:	נ ב מיתה		CTT(	سات ر ر	ነ ነ	Cac	I Tac	ארי אמרי	י דע	רדנידור V	מ מדרבר		ACAA	LLC. r	ACA	GAA	GCT(	STGA
, 721	C102	2361	مومد	9110	. 100	AIC.	CAC.	170	~~~										
+3	s i	? :	s v	L	P	P	s	A	L	P	L	D	L	L	N	N	A	I	T
781	GTC	TT	CAGT	GCT	ACCC	CCT	TCA	GCT	TTG	2 (	CACTG	GACC	T	GCTT	AAT.	AAT	GCT	ATA	ACTG
+3	A I	? :	s T	L	Ξ	D	L	I	R	Y	L	E	P	E	R	W	Q	L	D
841	CCT	rta(	STAC	CTTC	<b>GAA</b>	GAC	CIT	ATT	CGA:	r s	TCTT	GAAC	C	AGAG	AGA	TGG	CAG	TTG	GACT
_					_	_	_		_			_	••		_	••	_	_	<b>n</b>
د+ 901	L I		) L	Y		CC.	T	W TCC	Q CDA		T L	יים מפרז		CCCT	ملململ ت	ىنىڭ ۸	alana. E	ದರು.	2233
+3 961	K S	5 1	a v	v	D	L	N	L	L	T	E	E	kı	R	L	Y	s	С	T
961	AATO	CAC	GAGT	GGT	GAT	CTG	AAC	CTT	CTA	A (	CAGAG	GAÇ	; ;;	AAGA	TTA	TAC	AGC	TGC	ACAC
												<b>/</b> \							
+3	P I	2 1	N F	s	v	s	I	R	E	E	. L	K	R	T	D	T	I	F	W
1021	CTC	JTA.	ACTT	CTC	AGTG	TCC	ATA	AGG	GAA	G A	<b>ACTA</b>	aagi	\G	AACC	GAT	ACC	ATT	TIC	TGGC
			•													_			_
+3	P	3 (	C L	L	V	K	R	С	G	Ğ	N	C	A	С	C	L	H	N	C
1081	CAG	STT	GTCT	CCT	GTT	'AAA	CGC	TGT	GGT	G (	GAAC	TGT	ЭC	CTGT	IGT	CTC	CAC	AAT	IGCA
				_			_	7,0	**	-	. •	¥	v	Ľ	= :	v	7	^	•
+3	א איי	5 ( 33 T/	C Q	240	ρ <del>/ -</del> - ∀	.CC.	300	. A. T	المامات م	ν. Υ	K Taba	774. 7	47	CCAC	. = 242	Стс	باب ب	יראני. ע	TTG2
T141	MIG	JAN T	ar CA	AIG.	1616	· CCA	MGC	****	المنات	, ,	- 1 6464		• •			~~			4/1

FIG. 30 (CONTINUED 2).

+3 H E E C D C V C R G S T G G

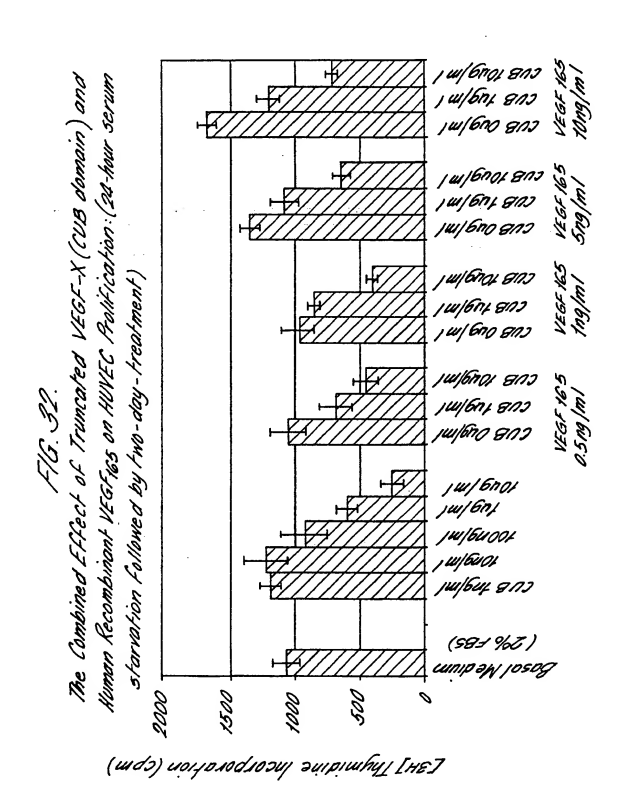
+3 R P K T G V R G L H K S L T D V A L E H
1201 GACCAAAGAC CGGTGTCAGG GGATTGCACA AATCACTCAC CGACGTGGCC CTGGAGCACC

1261 ATGAGGAGTG TGACTGTGTG TGCAGAGGGA GCACAGGAGG ATAGCCGCAT CACCACCAGC 1321 AGCTCTTGCC CAGAGCTGTG CAGTGCAGTG GCTGATTCTA TTAGAGAACG TATGCGTTAT 1381 CTCCATCCTT AAICTCAGTT GTTTGCTTCA AGGACCTTTC ATCTTCAGGA TTTACAGTGC 1441 ATTCTGAAAG AGGAGACATC AAACAGAATT AGGAGTTGTG CAACAGCTCT TTTGAGAGGA 1501 GGCCTAAAGG ACAGGAGAAA AGGTCTTCAA TCGTGGAAAG AAAATTAAAT GTTGTATTAA 1561 ATAGATCACC AGCTAGTTTC AGAGTTACCA TGTACGTATT CCACTAGCTG GGTTCTGTAT 1621 TTCAGTTCTT TCGATACGGC TTAGGGTAAT GTCAGTACAG GAAAAAAACT GTGCAAGTGA 1681 GCACCTGATT CCGTTGCCTT GCTTAACTCT AAAGCTCCAT GTCCTGGGCC TAAAATCGTA 1741 TAAAATCTGG ATTTTTTTT TTTTTTTTG CTCATATTCA CATATGTAAA CCAGAACATT 1801 CTATGTACTA CAAACCTGGT TTTTAAAAAG GAACTATGTT GCTATGAATT AAACTTGTGT 1861 CATGCTGATA GGACAGACTG GATTTTCAT ATTTCTTATT AAAATTTCTG CCATTTAGAA 1921 GAAGAGAACT ACATTCATGG TTTGGAAGAG ATAAACCTGA AAAGAAGAGT GGCCTTATCT 1981 TCACTTTATC GATAAGTCAG TTTATTTGTT TCATTGTGTA CATTTTTATA TTCTCCTTTT 2041 GACATTATAA CTGFTGGCTT TTCTAATCTT GTTAAATATA TCTATTTTTA CCAAAGGTAT 2101 TTAATATTCT TTTTTATGAC AACTTAGATC AACTATTTTT AGCTTGGTAA ATTTTTCTAA 2161 ACACAATTGT TATAGCCAGA GGAACAAAGA TGATATAAAA TATTGTTGCT CTGACAAAAA 2221 TACATGTATT TCATTCTCGT ATGGTGCTAG AGTTAGATTA ATCTGCATTT TAAAAAACTG 2281 AATTGGAATA GAATTGGTAA GTTGCAAAGA CTTTTTGAAA ATAATTAAAT TATCATATCT 2341 TCCATTCCTG TTATTGGAGA TGAAAATAAA AAGCAACTTA TGAAAGTAGA CATTCAGATC 2401 CAGCCATTAC TAACCTATTC CTTTTTTGGG GAAATCTGAG CCTAGCTCAG AAAAACATAA 2521 CACATCCTAT TIATTGTGAT GTTGTGGTTT TATTATCTTA AACTCTGTTC CATACACTTG 2581 TATAAATACA TGGATATTTT TATGTACAGA AGTATGTCTC TTAACCAGTT CACTTATTGT

2641 ACCTGGAAGG GCGAATTCTG CAGATATC

14/6nOx JW/6nf The Effect of FI-VEGF-X on HUVEC Poliferotion. j UI | 6U00F 14/6401 141641 100 6d 001 X-1931 (24-hour serum storvation followed by 14/640x 14/649 one day-treatment, jui/6uj שן 6009 | 1931 - 1931 בחירוכו (סחינס) MUIDAM JOSOG 1500 1000 (mds) suibinphi [HE]

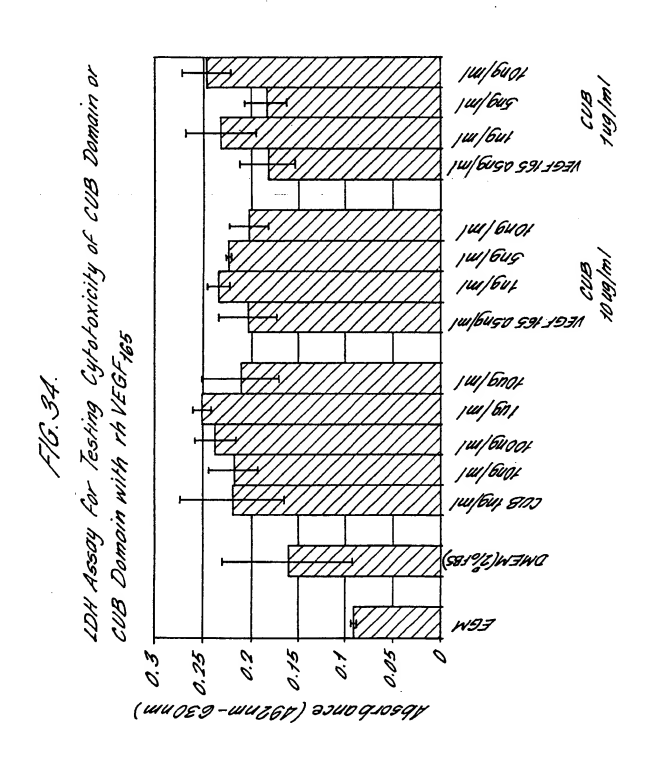
SUBSTITUTE SHEET (RULE 26)

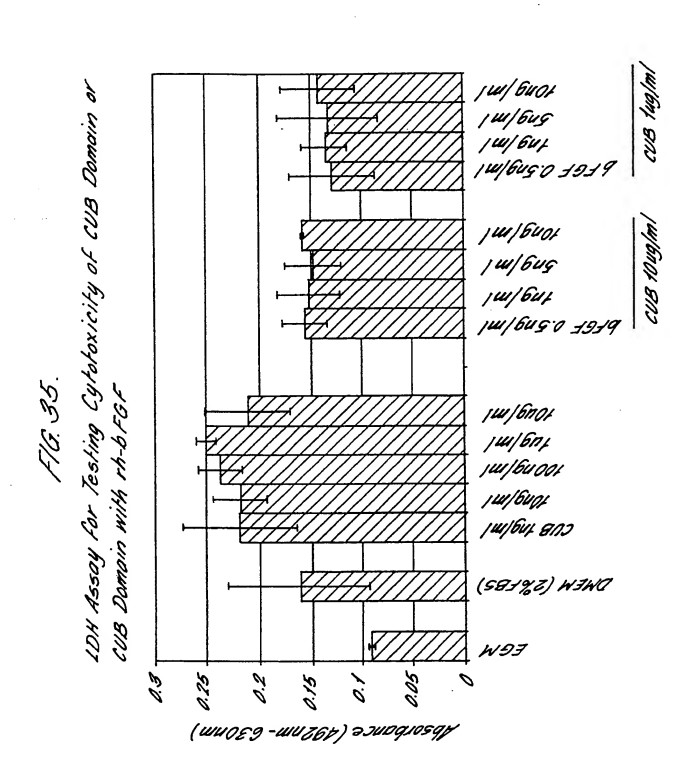


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F1G. 33. The Combined Effect of CUB Domain and Human Recombinant ruf6n0+ jul/6nf bFGF on HUVEC Poliferation: (24-hour serum starvation culbno ans jw/6not CNB 008/111 ju/6not jui/6nf 14/6no 8no followed by two-day-treatment) jw/6nof |w/bn/ |w/bn/ 8119 JUI/BNOF jui/6nj W/60001 141/6401 JUI/6UJ 811) (581%7) Bosol Medium 2000 2500 3000 3500 (mas) saidinital [HE]

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Applicant's oragent's file reference B0192/7011W0

131

International application No. PCT/US99/30503

### INDICATIONS RELATING TO DEPOSITED MICROORGANISM OR OTHER BIOLOGICAL MATERIAL

(PCT Rule 13bis)

A. The indications made below relate to the deposited microorgal on page $\begin{bmatrix} 21 \\ \end{bmatrix}$ . line $\begin{bmatrix} 15-16 \\ \end{bmatrix}$	nism or other biological material referred to in the description
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet
Name of depositary institution BELGIAN COORDINATED COLLECTIONS OF MI LABORATORIUM VOOR MOLECULAIRE BIOLOGI	
Address of depositary institution (including postal code and count Universiteit Gent K.L. Ledeganckstraat 35 B-9000 Gent, Belgium	יניד <i>ו</i>
Date of deposit	Accession Number
20 December 1999 (20.12.99)	LMBP 3991
C. ADDITIONAL INDICATIONS (leave blank if not applicable	(e) This information is continued on an additional sheet
D. DESIGNATED STATES FOR WHICH INDICATIONS A	
The indications listed below will be submitted to the International I Number of Deposit")	Bureau later (specify the general nature of the indications e.g., "Accession
For receiving Office use only	For International Bureau use only
This sheet was received with the international application	This sheet was received by the International Bureau on:  19 APRIL 2000 (19.84.60)
Authorized officer	Authorized officer Ellen Moyse

	ludapest Treaty on	the International Recognition of the Deposit of Microorganisms for the Purposes of Patent Procedure
lr	Receipt in the sternational Deposit	case of an original deposit issued pursuant to Rule 7.1 by the tary Authority BCCM <sup>TM</sup> /LMBP identified at the bottom of next page
		International Form BCCM™/LMBP/BP/4/99-23
To:	Name of the dep	ositor: Janssen Pharmaceutica N.V.
	Address	: Turnhoutseweg 30 B-2340 Beerse Belgium
ı.	Identification of	the microorganism:
	I.1 Identifica	ation reference given by the depositor:
	VEGF-X	CCUB PET22b
	I.2 Accessio	on number given by the International Depositary Authority:

# BELGIAN COORDINATED COLLECTIONS OF MICROORGANISMS - BCCM $^{\text{\tiny TM}}$ LMBP-COLLECTION

Page 2 of Form BCCM<sup>TM</sup>/LMBP/BP/4/99-23 Receipt in the case of an original deposit

II.	Scientific description and/or proposed taxono	mic designat	ion								
	The microorganism identified under I above w	as accompa	nied by	<b>/</b> :							
		(mark wi	th a cro	oss the applicable l	box(es))						
	<ul> <li>a scientific description</li> </ul>	yes	$\boxtimes$	no 🗌							
	<ul> <li>a proposed taxonomic designation</li> </ul>	yes		no 🛛							
III.	Receipt and acceptance										
	This International Depositary Authority accept above, which was received by it on (date of o	s the microo riginal depos	rganisı it) : D	m identified unde ecember 20, 199	er I 99						
IV.	International Depositary Authority										
	Belgian Coordinated Collections of Microorgan Laboratorium voor Moleculaire Biologie - Plasm Universiteit Gent K.L. Ledeganckstraat 35 B-9000 Gent, Belgium			P)							
	Signature(s) of person(s) having the power to a Authority or of authorized official(s):	represent the	e Interr	national Deposita	ary						
	-		anh.	Luke							

Martine Vanhoucke BCCM/LMBP curator

Date : January 12, 2000

On that date, the said microorganism was: (mark the applicable box with a cross)

図

viable

no longer viable

## BELGIAN COORDINATED COLLECTIONS OF MICROORGANISMS - $\mathsf{BCCM^{TM}}$ LMBP-COLLECTION

Page 2 of Form BCCM<sup>™</sup>/LMBP/BP/9/99-23 Viability statement

(Fill in if negative).	information	has	been	requested	and	if	the	results	of	the	test
		. <u>.</u>			<del></del>					<del></del> .	
		-									

V. International Depositary Authority

Belgian Coordinated Collections of Microorganisms (BCCM<sup>™</sup>)
Laboratorium voor Moleculaire Biologie - Plasmidencollectie (LMBP)
Universiteit Gent
K.L. Ledeganckstraat 35
B-9000 Gent, Belgium

Signature(s) of person(s) having the power to represent the International Depositary Authority or of authorized official(s):

Date : January 12, 2000

Martine Vanhoucke BCCM/LMBP curator